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Zur Erklärung der Zweibuchstaben-Codes und der anderen Abkürzungen wird auf die Erklärungen ("Guidance Notes on Codes and Abbreviations") am Anfang jeder regulären Ausgabe der PCT-Gazette verwiesen.

(54) Title: METHOD FOR DETERMINING HOMEOSTASIS OF THE SKIN

(54) Bezeichnung: VERFAHREN ZUR BESTIMMUNG DER HOMEOSTASE DER HAUT

(57) Abstract: The invention relates to a method for determining homeostasis of the skin in the skin of humans or animals in vitro. The invention also relates to test kits and biochips for determining homeostasis of the skin, in addition to the use of proteins, mRNA molecules or fragments of proteins or mRNA molecules as markers for homeostasis of the skin. The invention further relates to a test method for determining the efficacy of cosmetic or pharmaceutical active substances in order to maintain or promote homeostasis of the skin or in order to treat pathological conditions of the skin, as well as a screening method for the identification of cosmetic or pharmaceutical active substances for the maintenance or promotion of homeostasis of the skin or in order to treat pathological conditions of the skin and to a method for producing a cosmetic or pharmaceutical preparation in order to maintain or promote homeostasis of the skin or in order to treat pathological conditions of the skin.

(57) Zusammenfassung: Die vorliegende Erfindung betrifft ein Verfahren zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, Test-Kits und Biochips zur Bestimmung der Homeostase der haut sowie die Verwendung von Proteinen, mRNA-Molekülen oder Fragmenten von Proteinen oder mRNA-Molekülen als Marker für die Homeostase der Haut; ferner ein Testverfahren zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, sowie ein Screening-Verfahren zur Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut und ein Verfahren zur Herstellung einer kosmetischen oder pharmazeutischen Zubereitung zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut.

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#### Verfahren zur Bestimmung der Homeostase der Haut

Die vorliegende Erfindung betrifft ein Verfahren zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, Test-Kits und Biochips zur Bestimmung der Homeostase der Haut sowie die Verwendung von Proteinen, mRNA-Molekülen oder Fragmenten von Proteinen oder mRNA-Molekülen als Marker für die Homeostase der Haut; ferner ein Testverfahren zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, sowie ein Screening-Verfahren zur Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut und ein Verfahren zur Herstellung einer kosmetischen oder pharmazeutischen Zubereitung zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut bzw. zur Behandlung pathologischer Zustände der Haut.

Die Entwicklung eukaryotischen Lebens beginnt, abgesehen von der vegetativen Vermehrung, mit der Fusion zweier Gameten. Es entsteht eine Zygote, die der Ursprung einer jeden Zelle eines Eukaryoten ist. Die räumlich und zeitlich geordnete Differenzierung der Tochterzellen einer Zygote ist entscheidend für die Ontogenese eines vielzelligen Organismus, Sie führt zu verschiedensten Zelltypen, die sich in ihrer Morphologie und in ihrer Funktion unterscheiden. Vergleicht man beim Menschen z.B. eine Nervenzelle mit einer Zelle der Epidermis, so sind die Zellen sehr unterschiedlich, obwohl beide den gleichen Ursprung und das gleiche Genom haben. Die Differenzierung von Zellen geht mit Veränderung von Genexpressionsmustern einher. Im differenzierten Zustand exprimieren Zellen die für sie typischen Gene. Welche Gene dabei eine Rolle für die Morphologien und Funktionen z.B. der Hautzellen spielen ist bis heute

weitgehend unklar. Die geordnete Regulation der Genexpression in der Haut ist für die Aufrechterhaltung der Homeostase des Organs von entscheidender Bedeutung. Jede lebende Zelle ist in der Lage auf Signale ihrer Umwelt zu reagieren. Die Reaktionen der Zellen werden durch eine geordnete Regulation der Genexpression realisiert, sodaß der Metabolismus von Zellen nicht statisch sondern sehr dynamisch ist.

Die Expression der Gene in differenzierten Zellen der Haut ist nicht statisch sondern sehr dynamisch. Extrazelluläre Stimuli wirken über zum Teil komplexe Signaltransduktionskaskaden auf die Transkription lebender Zellen. Die Regulation der Transkription als Antwort auf extrazelluläre Signale wird als Stimulus-Transkriptions-Kopplung bezeichnet. Die Beeinflussung dieses empfindlichen Regulationsmechanismus kann zur Störung der Homeostase der Haut und möglicherweise zur Entstehung und Manifestation pathogener Zustände der Haut führen.

Das menschliche Genom umfasst nach jüngsten Schätzungen ca. 140 000 Gene. Von diesem immensen Informationsangebot verwendet jede Zelle jedoch lediglich einen kleinen, für sie spezifischen Teil für die Synthese von Proteinen, der sich im Genexpressionsmuster wiederspiegelt. Welche Gene insbesondere in der Haut eine Rolle spielen ist bisher weitgehend unklar.

Die Haut ist das größte Organ des menschlichen Körpers. Sie ist ein sehr komplex aufgebautes Organ, welches aus einer Vielzahl verschiedener Zelltypen besteht und die Grenzfläche des Körpers zur Umwelt bildet. Diese Tatsache verdeutlicht, dass die Zellen der Haut in besonderem Maße exogenen Signalen der Umwelt, physikalischer und chemischer Natur ausgesetzt. Für das Verständnis von Hautreaktionen auf exogene Stimuli ist die Analyse der Genexpression in der Haut von entscheidender Bedeutung.

Ein entscheidendes Merkmal der Haut ist, dass mit zunehmendem Alter, unter dem Einfluss hautschädigender Stimuli oder bei pathologischen Zuständen der Haut die Zellen ihre Fähigkeit verlieren die Homeostase des Organs aufrecht zu erhalten. Welche molekularen Mechanismen dieser Entwicklung zugrunde liegen ist bislang weitgehend unklar. Die Identifikation neuer hautspezifischer Marker ermöglicht, den komplexen Zustand der Homöostase, die Entstehung und Manifestation hautpathogener Zustände zu begreifen. Nur mit diesem Wissen können neue Konzepte für Produkte zur Hautbehandlung entwickelt werden.

Jeder Zelltyp der Haut exprimiert ca. 15.000 verschiedene Gene und synthetisiert daraus entsprechend viele Proteine. Welche Gene davon für die Homeostase der Haut eine Rolle spielen oder an pathogenen Prozessen beteiligt sind ist bisher jedoch weitgehend unklar.

Die Haut besteht aus mehreren verschiedenen Zelltypen (Fibroblasten, Keratinozyten in verschiedenen Differenzierungszuständen, Melanozyten, Merkelzellen, Langerhanszellen Haarfollikelzellen, Schweisdrüsenzellen etc.), sodass die Komplexität in der Haut exprimierter Gene sehr groß ist. Es ist bisher nicht möglich gewesen, diese immense Komplexität zu beschreiben. Ebenso wenig war es bisher möglich aus dieser Komplexität die Gene zu identifizieren, die exklusiv bzw. besonders stark in der Haut exprimiert werden.

In lebenden Zellen kommen mRNA-Moleküle in Konzentrationen zwischen einigen wenigen und mehreren hundert Kopien vor. Die schwach exprimierten Gene sind bisherigen Analysen nicht oder nur sehr schwer zugänglich gewesen. Diese Moleküle können aber durchaus eine entscheidende Rolle für die Homeostase der Haut spielen oder an der Entstehung bzw. Manifestation pathogener Prozesse in der Haut beteiligt sein.

Die Gesamtheit aller mRNA-Moleküle, die von einer Zelle oder einem Gewebe zu einem bestimmten Zeitpunkt synthetisiert werden, bezeichnet man als "Transkriptom". Bis heute ist es nicht möglich gewesen das komplette Transkriptom, also die Gesamtheit aller transkribierten Gene, der humanen Haut zu beschrieben.

Die Analyse der Genexpression ist zwar mit der Quantifizierung spezifischer mRNA-Moleküle möglich (z.B. Northern-Blot, RNase-Schutzexperimente). Mit diesen Techniken können jedoch nur eine relativ begrenzte Anzahl an Genen gemessen werden.

Es besteht daher ein Bedarf an der Identifikation möglichst vieler, vorzugsweise aller, in menschlicher oder tierischer Haut aktiven Gene.

Aufgabe der vorliegenden Erfindung ist es daher, einen möglichst großen Teil der in menschlicher oder tierischer Haut exprimierten Gene zu identifizieren; ferner, die für die Homeostase der Haut bedeutsamen Gene zu identifizieren. Außerdem sollen mittels der identifizierten Gene, Verfahren zur Bestimmung der Homeostase der Haut bereitgestellt werden.

Diese erste Aufgabe wird erfindungsgemäß gelöst durch ein Verfahren (1) zur Identifizierung der in Haut exprimierten Gene bei Menschen oder Tieren in vitro, das dadurch gekennzeichnet ist, daß man

- a) ein Gemisch von in menschlicher oder tierischer Haut exprimierten, d. h. transkribierten genetisch codierten Faktoren, also von mRNA-Molekülen oder Fragmenten von mRNA-Molekülen aus menschlicher oder tierischer Haut gewinnt und
- b) das in a) gewonnenen Gemisch einer Seriellen Analyse der Genexpression (SAGE) unterwirft, und dadurch die in menschlicher oder tierischer Haut exprimierten Gene identifiziert und ihre Expression quantifiziert.

Die zweite Aufgabe wird erfindungsgemäß gelöst durch ein Verfahren (2) zur Identifizierung der für die Homeostase der Haut bedeutsamen Gene bei Menschen oder Tieren in vitro, das dadurch gekennzeichnet ist, daß man

a) ein Gemisch von in menschlicher oder tierischer Haut exprimierten, d. h. transkribierten genetisch codierten Faktoren, also von mRNA-Molekülen oder

Fragmenten von mRNA-Molekülen aus menschlicher oder tierischer Haut gewinnt,

- b) das in a) gewonnenen Gemisch einer Seriellen Analyse der Genexpression (SAGE) unterwirft, und dadurch die in menschlicher oder tierischer Haut exprimierten Gene identifiziert und ihre Expression quantifiziert und
- c) die Analysergebnisse aus b) mit Expressionsmustern anderer Gewebe vergleicht und so die Gene identifiziert, die in Haut und anderen Geweben unterschiedlich stark (differentiell) exprimiert werden.

Expressionsmuster anderer Gewebe sind beispielsweise in den Datenbanken des Cancer Genome Anatomy Project (CGAP) im Internet unter folgender Adresse zugänglich: http://cgap.nci.nih.gov/

Zur Erfassung des Transkriptoms der Haut wurde die Technik der "Seriellen Analyse der Genexpression" (SAGE™) eingesetzt. Diese Technik erlaubt gleichzeitig die Identifikation und Quantifizierung aller in der Haut exprimierten Gene. Der Vergleich des Transkriptoms der Haut, mit dem Transkriptom anderer Gewebe lässt die Unterscheidung zwischen relevanten und nicht relevanten Genen für die Homeostase der Haut zu.

Für die SAGE™-Analyse wurde humane Haut von gesunden weiblichen Spendern verwendet. Die Durchführung der SAGE™-Analyse erfolgte wie in der EP-A-0 761 822 und bei Velculescu, V.E. et al., 1995 Science 270, 484-487, beschrieben und führte zur Identifikation der in Haut aktiven Gene.

Diese Gene sind dazu geeignet die Homeostase der Haut zu bestimmen oder pathologische Prozesse oder Zustände zu detektieren.

Die Tabelle 6 enthält eine detaillierte Auflistung der mit Hilfe des erfindungsgemäßen Verfahrens (1) ermittelten, in menschlicher Haut aktiven Gene unter Angabe

- einer laufenden Ordnungsnummer in Spalte 1,
- der verwendeten Tag-Sequenz in Spalte 2,

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- der ermittelten relativen Expressionsfrequenz in Haut in Spalte 3,
- der Signifikanz in Spalte 4,
- der UniGene-Accession-Number in Spalte 5 und
- einer Kurzbeschreibung des Gens bzw. Genproduktes in Spalte 6.

Die Tabellen 1 bis 5 enthalten eine detaillierte Auflistung der mit Hilfe des erfindungsgemäßen Verfahrens (2) ermittelten, in Haut und in anderen Geweben differentiell exprimierten Gene unter Angabe

- einer laufenden Ordnungsnummer in Spalte 1,
- der verwendeten Tag-Sequenz in Spalte 2,
- der ermittelten relativen Expressionsfrequenz im CGAP (Cancer Genome Anatomy Project) in Spalte 3,
- der ermittelten relativen Expressionsfrequenz in Haut in Spalte 4,
- des Quotienten der Frequenzen (aus Spalte 3 und Spalte 4) in Spalte 5,
- der Signifikanz in Spalte 6,
- der UniGene-Accession-Number in Spalte 7 und
- einer Kurzbeschreibung des Gens bzw. Genproduktes in Spalte 8.

Der Quotient in Spalte 5 gibt die Stärke der differentiellen Expression an, d. h., um welchen Faktor das jeweilige Gen in Haut stärker exprimiert wird, als in anderen Geweben.

#### In Tabelle 7 sind unter Angabe

- einer laufenden Ordnungsnummer in Spalte 1,
- der verwendeten Tag-Sequenz in Spalte 2,
- der ermittelten relativen Expressionsfrequenz im CGAP (Cancer Genome Anatomy Project) in Spalte 3,
- der ermittelten relativen Expressionsfrequenz in Haut in Spalte 4,
- des Quotienten der Frequenzen (aus Spalte 3 und Spalte 4) in Spalte 5,
- der UniGene-Accession-Number in Spalte 6 und
- einer Kurzbeschreibung des Gens bzw. Genproduktes in Spalte 7,

Gene aufgelistet, die zwischen 13,33- und 211,11-fach differentiell exprimiert sind. Die Zuordnung der Tags zu den Genen, die durch Ihre UniGene-Accession-Number in Spalte 6 definiert werden, erfolgte durch manuelle Annotation.

Zur Annotation wurden folgende Datenbanken verwendet:

- 1. Unigene Version vom 30.10.01 mit folgenden Datenbankeinträgen:
  - a. der bekannten Gene aus Genbank (Stand: 12.10.01)
  - b. der EST's aus dbEST (Stand: 19.10.01)
- 2. mRNA Version released am 17.10.01

Die Datenbanken wurden vom NCBI heruntergeladen, für eine lokale Version des BLAST-Programmes (ebenfalls NCBI) formatiert und mit den in der SAGE-Analyse detektierten Tags auf identische Hits verglichen.

Die gefundenen Gene/Klone wurden auf Redundanz geprüft und wie nachfolgend aufgeführt nachbearbeitet:

- Tag-Sequenzen mit mehreren unterschiedlichen Treffern: Bewertung als nicht annotierbar.
- 2. Tag-Sequenzen mit doppelten oder mehreren identischen Treffern: Eliminierung der Treffer, die am weitesten vom Poly-A-Tail entfernt lagen.

Zunächst wurden die Ergebnisse aus der Unigene-Datenbank ausgewertet und dann mit den Ergebnissen aus der mRNA-Datenbank abgeglichen. Letztere tauchen in der Tabelle 7 nicht auf, da sie auch über die Unigene-Einträge abrufbar sind.

Alle in der Ergebnistabelle aufgeführten Links wurden auf der im folgenden dokumentierten Datenbasis des 30.10.2001 (Unigene-Datenbankrelease: UniGene Build #143) überprüft:

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#### Sequences Included in UniGene

Known genes are from GenBank (Oct 12, 2001) ESTs are from dbEST through 19-Oct-2001

69367 mRNAs + gene CDSs

1147828 EST, 3'reads

1196006 EST, 5'reads

+ 598081 EST, other/unknown

3011282 total sequences in clusters

### Final Number of Clusters (sets)

\_\_\_\_\_\_

96332 sets total

20516 sets contain at least one known gene

95171 sets contain at least one EST

19355 sets contain both genes and ESTs

#### Release Notes

Unter ihrer UniGene-Accession-Number sind die jeweiligen Gene bzw. Genprodukte in der Datenbank des National Center for Biotechnology Information (NCBI) offenbart. Diese Datenbank ist im Internet unter folgender Adresse zugänglich: <a href="http://www.ncbi.nlm.nih.gov/">http://www.ncbi.nlm.nih.gov/</a>.

Die Gene bzw. Genprodukte sind außerdem unter den Internet-Adressen <a href="http://www.ncbi.nlm.nih.gov/UniGene/Hs.Home.html">http://www.ncbi.nlm.nih.gov/UniGene/Hs.Home.html</a> oder <a href="http://www.ncbi.nlm.nih.gov/genome/guide">http://www.ncbi.nlm.nih.gov/genome/guide</a> direkt zugänglich.

Die Daten des Cancer Genome Anatomy Project sind im Internet unter folgender Adresse zugänglich: <a href="http://cgap.nci.nih.gov/">http://cgap.nci.nih.gov/</a>

In Tabelle 1 sind alle Gene aufgelistet, die mindestens 2-fach und weniger als 5fach differentiell exprimiert sind.

In Tabelle 2 sind alle Gene aufgelistet, die mindestens 5-fach und weniger als 10-fach differentiell exprimiert sind.

In Tabelle 3 sind alle Gene aufgelistet, die mindestens 10-fach und weniger als 20-fach differentiell exprimiert sind.

In Tabelle 4 sind alle Gene aufgelistet, die mindestens 20-fach und weniger als 100-fach differentiell exprimiert sind.

In Tabelle 5 sind alle Gene aufgelistet, die mindestens 100-fach differentiell exprimiert sind.

Die dritte der vorliegenden Erfindung zugrundeliegende Aufgabe wird erfindungsgemäß gelöst durch ein Verfahren (3) zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren, insbesondere bei Frauen, in vitro, das dadurch gekennzeichnet ist, daß man

- a) ein Gemisch von Proteinen, mRNA-Molekülen oder Fragmenten von Proteinen oder mRNA-Molekülen aus menschlicher oder tierischer Haut gewinnt,
- b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die mittels Serieller Analyse der Genexpression (SAGE) als in Haut und anderen Geweben unterschiedlich stark (differentiell) exprimiert identifiziert werden,
- c) die Untersuchungsergebnisse aus b) mit den mittels Serieller Analyse der Genexpression (SAGE) identifizierten Expressionsmustern vergleicht und

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d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut stärker exprimiert werden als in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben stärker exprimiert werden als in Haut.

Es kann in Schritt b) des Verfahrens zur Bestimmung der Homeostase der Haut ausreichend sein, das gewonnene Gemisch auf das Vorhandensein von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen zu untersuchen, die mittels Serieller Analyse der Genexpression (SAGE) als in Haut und anderen Geweben differentiell exprimiert identifiziert werden, wenn diese ausschließlich in Haut oder ausschließlich in anderen Geweben exprimiert werden. In allen anderen Fällen muß in Schritt b) auch die Menge der differentiell exprimierten Moleküle untersucht werden, d. h., die Expression muß quantifiziert werden.

In Schritt d) des Verfahrens zur Bestimmung der Homeostase der Haut wird das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zugeordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut stärker exprimiert werden als in anderen Geweben, d. h., daß das Gemisch entweder mehr unterschiedliche typischerweise in Haut exprimierte Verbindungen enthält, als solche, die typischerweise in anderen Geweben exprimiert werden Differenzierung), oder mehr Kopien von typischerweise in Haut exprimierten Verbindungen enthält, als typischerweise in anderen Geweben vorhanden sind (quantitative Differenzierung). Für die Zuordnung zu kranker bzw. in gestörter Homeostase befindlicher Haut wird in komplementärer Weise verfahren.

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Eine bevorzugte Ausführungsform des erfindungsgemäßen Verfahrens zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 1 bis 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens doppelt so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens doppelt so stark exprimiert werden wie in Haut.

Eine weitere bevorzugte Ausführungsform des erfindungsgemäßen Verfahrens zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 2 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 2 bis 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens

5-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 5-fach so stark exprimiert werden wie in Haut.

Eine weitere bevorzugte Ausführungsform des erfindungsgemäßen Verfahrens zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 3 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 3 bis 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 10-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 10-fach so stark exprimiert werden wie in Haut.

Eine weitere bevorzugte Ausführungsform des erfindungsgemäßen Verfahrens zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 4 und 5 in Spalte 7 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 4 und 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 20-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 20-fach so stark exprimiert werden wie in Haut.

Eine weitere bevorzugte Ausführungsform des erfindungsgemäßen Verfahrens zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in Tabelle 5 in Spalte 7 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in Tabelle 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 100-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 100-fach so stark exprimiert werden wie in Haut.

Man kann den Zustand der Haut auch dadurch beschreiben, daß mehrere Marker (Expressionprodukte der für die Homeostase der Haut bedeutsamen Gene)

quantifiziert werden, die dann untereinander in einem bestimmten Verhältnis aktiv sein müssen, um in Homeostase befindliche Haut zu repräsentieren. Alle Abweichungen hiervon deuten darauf hin, daß die untersuchte Haut sich nicht in Homeostase befindet.

Ein weiterer Gegenstand der vorliegenden Erfindung ist daher ein Verfahren (4) zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren, insbesondere bei Frauen, in vitro, das dadurch gekennzeichnet ist, daß man

- a) ein Gemisch von Proteinen, mRNA-Molekülen oder Fragmenten von Proteinen oder mRNA-Molekülen aus menschlicher oder tierischer Haut gewinnt,
- b) in dem gewonnenen Gemisch mindestens zwei der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen quantifiziert, die mittels Verfahren (2) als für die Homeostase der Haut bedeutsam identifiziert werden,
- c) die Expressionsverhältnisse der mindestens zwei Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen zueinander bestimmt,
- d) die Expressionsverhältnisse aus c) mit den Expressionsverhältnissen vergleicht, die für die in b) quantifizierten Moleküle typischerweise in homeostatischer Haut vorliegen, insbesondere mit den Expressionsverhältnissen, die sich aus Tabelle 6, Spalte 3 bzw. aus den Tabellen 1 bis 5, Spalte 4 ergeben, und
- e) das in a) gewonnene Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn die Expressionsverhältnisse der untersuchten Haut den Expressionsverhältnissen in Homeostase befindlicher Haut entsprechen, oder das in a) gewonnene Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn die Expressionsverhältnisse der untersuchten Haut von den Expressionsverhältnissen in Homeostase befindlicher Haut abweichen.

Vorzugsweise gewinnt man in Schritt a) der erfindungsgemäßen Verfahren zur Bestimmung der Homeostase der Haut das Gemisch aus einer Hautprobe, insbesondere aus einer Vollhautprobe oder aus einer Epidermisprobe. Hierbei eröffnet die Vollhautprobe umfassendere Vergleichsmöglichkeiten mit den

gleichfalls aus Vollhaut gewonnenen SAGE-Libraries. Die Epidermisprobe ist hingegen leichter zu gewinnen, beispielsweise durch Aufbringen eines Klebebandes auf die Haut und Abreißen desselben, wie in der WO 00/10579 beschrieben, auf die hiermit in vollem Umfang Bezug genommen wird.

In einer weiteren Ausführungsform der erfindungsgemäßen Verfahren zur Bestimmung der Homeostase der Haut gewinnt man in Schritt a)

das Gemisch mittels Mikrodialyse. Die Technik der Mikrodialyse wird beispielsweise in "Microdialysis: A method for measurement of local tissue metabolism", Nielsen PS, Winge K, Petersen LM; Ugeskr Laeger 1999 Mar 22 161:12 1735-8; sowie in "Cutaneous microdialysis for human in vivo dermal absorption studies", Anderson, C. et al.; Drugs Pharm. Sci., 1998, 91, 231-244; und auch im Internet unter http://www.microdialysis.se/techniqu.htm beschrieben, worauf hiermit in vollem Umfang Bezug genommen wird.

Bei der Anwendung der Mikrodialyse führt man typischerweise eine Sonde in die Haut ein und beginnt mit einer geeigneten Trägerlösung die Sonde langsam zu spülen. Nach dem Abklingen der akuten Reaktionen nach dem Einstich liefert die Mikrodialyse Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen, die im extrazellulären Raum vorkommen und die, beispielsweise durch Fraktionierung der Trägerflüssigkeit, dann in vitro isoliert und analysiert werden können. Die Mikrodialyse ist weniger invasiv, als die Entnahme einer Vollhautprobe; sie ist aber nachteiligerweise auf die Gewinnung im extrazelulären Raum vorkommender Verbindungen beschränkt.

Eine weitere bevorzugte Ausführungsform der erfindungsgemäßen Verfahren zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) in Verfahren (3) die Untersuchung auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine oder Proteinfragmente; bzw. in Verfahren (4) die Quantifizierung mindestens zweier Proteine oder Proteinfragmente, mittels einer Methode durchführt, die ausgewählt ist unter

Ein- oder zweidimensionaler Gelelektrophorese

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- Affinitätschromatographie
- Protein-Protein-Komplexierung in Lösung
- Massenspektrometrie, insbesondere Matrix Assistierter Laser Desorptions Ionisation (MALDI) und insbesondere
- Einsatz von Proteinchips, oder mittels geeigneter Kombinationen dieser Methoden.

Diese erfindungsgemäß einsetzbaren Methoden sind in dem Übersichtsartikel von Akhilesh Pandey und Matthias Mann: "Proteomics to study genes and genomes", Nature, Volume 405, Number 6788, 837 - 846 (2000), und den dort angegebenen Referenzen beschrieben, worauf hiermit in vollem Umfang Bezug genommen wird.

Die 2D-Gelelektrophorese, wird beispielsweise in L.D. Adams, Two-dimensional Gel Electrophoresis using the Isodalt System oder in L.D. Adams & S.R. Gallagher, Two-dimensional Gel Electrophoresis using the O'Farrell System; beide in Current Protocols in Molecular Biology (1997, Eds. F.M. Ausubel et al.), Unit 10.3.1 - 10.4.13; oder in 2-D Electrophoresis-Manual; T. Berkelman, T. Senstedt; Amersham Pharmacia Biotech, 1998 (Bestell-Nr. 80-6429-60), beschrieben.

Die massenspektrometrische Charakterisierung der **Proteine** oder Proteinfragmente erfolgt in der Fachwelt bekannter Weise, beispielsweise wie in den folgenden Literaturstellen beschrieben:

Methods in Molecular Biology, 1999; Vol 112; 2-D Proteome Analysis Protocols; Editor: A. J. Link; Humana Press; Totowa; New Jersey. Darin insbesondere: Courchesne, P. L. und Patterson, S. D.; S. 487-512.

Carr, S. A. und Annan, R. S.; 1997; in: Current Protocols in Molecular Biology; Editor: Ausubel, F. M. et al.; John Wiley and Sons, Inc. 10.2.1-10.21.27.

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Eine weitere bevorzugte Ausführungsform der erfindungsgemäßen Verfahren zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) in Verfahren (3) die Untersuchung auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der mRNA-Moleküle oder mRNA-Molekülfragmente; bzw. in Verfahren (4) die Quantifizierung mindestens zweier mRNA-Moleküle oder mRNA-Molekülfragmente mittels einer Methode durchführt, die ausgewählt ist unter

- i. Northern Blots,
- ii. Reverse Transkriptase Polymerasekettenreaktion (RT-PCR),
- iii. RNase-Schutzexperimente,
- iv. Dot-Blots,
- v. cDNA-Sequenzierung,
- vi. Klon-Hybridisierung,
- vii. Differential Display,
- viii. Subtraktive Hybridisierung,
- ix. cDNA-Fragment-Fingerprinting,
- x. Total Gene Expression Analysis (TOGA)
- xi. Serielle Analyse der Genexpression (SAGE) und insbesondere
- xii. Einsatz von Nukleinsäurechips,

oder mittels geeigneter Kombinationen dieser Methoden.

Diese erfindungsgemäß einsetzbaren Methoden sind in den Übersichtsartikeln von Akhilesh Pandey und Matthias Mann: "Proteomics to study genes and genomes", Nature, Volume 405, Number 6788, 837 - 846 (2000), und "Genomics, gene expression and DNA arrays", Nature, Volume 405, Number 6788, 827 - 836 (2000), und den dort angegebenen Referenzen beschrieben, worauf hiermit in vollem Umfang Bezug genommen wird.

Das TOGA-Verfahren ist in "J. Gregor Sutcliffe et al, TOGA: An automated parsing technology for analyzing expression of nearly all genes, Proceedings of the National Academy of Sciences of the United States of America (PNAS), Vol. 97, No. 5, pp. 1976-1981 (2000)" beschrieben, worauf hiermit vollumfänglich Bezug genommen wird.

Es können jedoch erfindungsgemäß auch andere dem Fachmann bekannte Methoden zur Untersuchung auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen eingesetzt werden.

Eine weitere bevorzugte Ausführungsform der erfindungsgemäßen Verfahren zur Bestimmung der Homeostase der Haut ist dadurch gekennzeichnet, daß man in Schritt b) auf das Vorhandensein und gegebenenfalls die Menge von 1 bis etwa 5000, bevorzugt 1 bis etwa 1000, insbesondere etwa 10 bis etwa 500, vorzugsweise etwa 10 bis etwa 250, besonders bevorzugt etwa 10 bis etwa 100 und ganz besonders bevorzugt etwa 10 bis etwa 50 der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden.

Ein weiterer Gegenstand der vorliegenden Erfindung ist ein Test-Kit zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, umfassend Mittel zur Durchführung der erfindungsgemäßen Verfahren zur Bestimmung der Homeostase der Haut.

Ein weiterer Gegenstand der vorliegenden Erfindung ist ein Biochip zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, umfassend

- i. einen festen, d. h. starren oder flexiblen Träger und
- ii. auf diesem immobilisierte Sonden, die zur spezifischen Bindung an mindestens eines der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen befähigt sind, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden.

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Bei einem BioChip handelt es sich um ein miniaturisiertes Funktionselement mit auf einer Oberfläche immobilisierten Molekülen, insbesondere Biomolekülen, die als spezifische Interaktionspartner dienen können.

Häufig weist die Struktur dieser Funktionselemente Reihen und Spalten auf; man spricht dann von Chip-"Arrays". Da tausende von biologischen bzw. biochemischen Funktionselementen auf einem Chip angeordnet sein können, müssen diese in der Regel mit mikrotechnischen Methoden angefertigt werden.

Als biologische und biochemische Funktionselemente kommen insbesondere in Frage: DNA, RNA, PNA, (bei Nukleinsäuren und ihren chemischen Derivaten können z. B. Einzelstränge, Triplex-Strukturen oder Kombinationen hiervon vorliegen), Saccharide, Peptide, Proteine (z. B. Antikörper, Antigene, Rezeptoren) und Derivate der kombinatorischen Chemie (z. B. organische Moleküle).

Im allgemeinen haben BioChips eine 2D-Basisfläche für das Beschichten mit biologisch oder biochemisch funktionellen Materialien. Die Basisflächen können beispielweise auch von Wänden einer oder mehrerer Kapillaren oder von Kanälen gebildet sein.

Zum Stand der Technik kann z. B. auf folgende Publikationen hingewiesen werden: Nature Genetics, Vol. 21, supplement (Gesamt), Jan. 1999 (BioChips); Nature Biotechnology, Vol. 16, S. 981-983, Okt. 1998 (BioChips); Trends in Biotechnology, Vol. 16, S. 301-306, Jul. 1998 (BioChips) sowie die bereits genannten Übersichtsartikel von Akhilesh Pandey und Matthias Mann: "Proteomics to study genes and genomes", Nature, Volume 405, Number 6788, 837 - 846 (2000), und "Genomics, gene expression and DNA arrays", Nature, Volume 405, Number 6788, 827 - 836 (2000), und die dort angegebenen Referenzen, worauf hiermit in vollem Umfang Bezug genommen wird.

Eine übersichtliche Darstellung der praktischen Anwendungsverfahren der DNA-Chiptechnologie liefern die Bücher "DNA Microarrays: A Practical Approach" (Editor: Mark Schena, 1999, Oxford University Press) und "Microarray Biochip Technology" (Editor: Mark Schena, 2000, Eaton Publishing), auf die hiermit in vollem Umfang Bezug genommen wird.

Die im Rahmen der vorliegenden Erfindung besonders bevorzugte DNA-Chiptechnologie beruht auf der Fähigkeit von Nukleinsäuren komplementäre Basenpaarungen einzugehen. Dieses als Hybridisierung bezeichnete technische Prinzip wird bereits seit Jahren bei der Southern-Blot- und Northern-Blot-Analyse eingesetzt. Im Vergleich zu diesen herkömmlichen Methoden, bei denen lediglich einige wenige Gene analysiert werden, gestattet es die DNA-Chiptechnologie einige hundert bis zu mehreren zehntausend Genen parallel zu untersuchen.

Ein DNA-Chip besteht im wesentlichen aus einem Trägermaterial (z.B. Glas oder Kunststoff), auf dem einzelsträngige, genspezifische Sonden in hoher Dichte an einer definierten Stelle (Spot) immobilisiert werden. Als problematisch wird dabei die Technik der Sonden-Applikation und die Chemie der Sonden-Immobilisierung eingeschätzt.

Nach dem derzeitigen Stand der Technik sind mehrere Wege der Sonden-Immobilisierung realisiert:

E.M. Southern (E.M. Southern et al. (1992), Nucleic Acid Research 20, 1679-1684 und E.M. Southern et al. (1997), Nucleic Acid Research 25, 1155-1161) beschreibt die Herstellung von Oligonukleotidanordnungen durch direkte Synthese an einer Glasoberfläche, die mit 3-Glycidoxypropyltrimethoxysilan und anschließend mit einem Glycol derivatisiert wurde.

Ein ähnliches Verfahren realisiert die *in situ* Synthese von Oligonukleotiden mittels einer photosensitiven, kombinatorischen Chemie, die mit photolithographischen Techniken verglichen werden kann (Pease, A.C. et al. (1994), Proc. Natl Acad Sci USA 91, 5022-5026).

Neben diesen auf der *in situ-*Synthese von Oligonukleotiden beruhenden Techniken können ebenso bereits vorhandene DNA-Moleküle an Oberflächen von Trägermaterial gebunden werden.

P.O. Brown (DeRisi et al. (1997), Science 278, 680-686) beschreibt die Immobilisierung von DNA an mit Polylysin beschichteten Glasoberflächen.

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Die Veröffentlichung von L.M. Smith (Guo, Z. et al. (1994), Nucleic Acid Research 22, 5456-5465) legt ein ähnliches Verfahren offen: Oligonukleotide, die eine 5'terminale Aminogruppe tragen, können an eine Glasoberfläche gebunden werden, die mit 3-Aminopropyltrimethoxysilan und anschließend mit 1,4-Phenyldiisothiocyanat behandelt wurde.

Die Applikation der DNA-Sonden auf einem Träger kann mit einem sogenannten "Pin-Spotter" erfolgen. Dazu tauchen dünne Metallnadeln mit z.B. einem Durchmesser von 250 µm, in Sondenlösungen ein und überführen anschließend das anhängende Probenmaterial mit definierten Volumina auf das Trägermaterial des DNA-Chips.

Bevorzugterweise erfolgt die Sondenapplikation jedoch mittels eines piezogesteuerten Nanodispensers, der ähnlich einem Tintenstrahldrucker, Sondenlösungen mit einem Volumen von 100 Picolitern kontaktfrei auf die Oberfläche des Trägermaterials aufbringt.

Die Immobilisierung der Sonden erfolgt z.B. wie in der EP-A-0 965 647 beschrieben: Die Generierung von DNA-Sonden erfolgt hierbei mittels PCR unter Verwendung eines sequenzspezifischen Primerpaares, wobei ein Primer am 5'-Ende modifiziert ist und einen Linker mit einer freien Aminogruppe trägt. Damit ist sichergestellt, dass ein definierter Strang der PCR-Produkte an einer Glasoberfläche gebunden werden kann, welche mit 3-Aminopropyltrimethoxysilan und anschließend mit 1,4-Phenyldiisothiocyanat behandelt wurde. genspezifischen **PCR-Produkte** sollen idealerweise eine definierte Nukleinsäuresequenz in einer Länge von 200-400 bp haben und nicht redundante Seguenzen beinhalten. Nach der Immobilisierung der PCR-Produkte über den derivatisierten Primer wird der Gegenstrang des PCR-Produkts durch eine Inkubation bei 96°C für 10 Min entfernt.

In einer für DNA-Chips typischen Anwendung wird mRNA aus zwei zu vergleichenden Zellpopulationen isoliert. Die isolierten mRNAs werden mittels

reverser Transkription unter Verwendung von z.B. fluoreszenzmarkierten Nukleotiden in cDNA umgewandelt. Dabei werden die zu vergleichenden Proben mit z.B. rot bzw. grün fluoreszierenden Nukleotiden markiert. Die cDNAs werden dann mit den auf dem DNA-Chip immobilisierten Gensonden hybridisiert und anschließend die gebundenen Fluoreszenzen quantifiziert.

Der erfindungsgemäße Biochip umfasst bevorzugt 1 bis etwa 5000, bevorzugtermaßen 1 bis etwa 1000, insbesondere etwa 10 bis etwa 500, vorzugsweise etwa 10 bis etwa 250, besonders bevorzugt etwa 10 bis etwa 100 und ganz besonders bevorzugt etwa 10 bis etwa 50 voneinander verschiedene Sonden. Die voneinander verschiedenen Sonden können jeweils in mehrfacher Kopie auf dem Chip vorhanden sein.

Der erfindungsgemäße Biochip umfasst bevorzugt Nukleinsäuresonden, insbesondere RNA- oder PNA-Sonden, besonders bevorzugt DNA-Sonden. Die Nukleinsäuresonden weisen bevorzugt eine Länge von etwa 10 bis etwa 1000, insbesondere etwa 10 bis etwa 800, vorzugsweise etwa 100 bis etwa 600, besonders bevorzugt etwa 200 bis etwa 400 Nukleotiden auf.

In einer weiteren bevorzugten Form umfasst der erfindungsgemäße Biochip Peptid- oder Proteinsonden, insbesondere Antikörper.

Ein weiterer Gegenstand der vorliegenden Erfindung ist die Verwendung der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden, als Marker für die Homeostase der Haut bei Menschen oder Tieren.

Ein weiterer Gegenstand der vorliegenden Erfindung ist ein Testverfahren zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand,

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Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom in vitro, dadurch gekennzeichnet, daß man

- a) den Hautstatus durch ein erfindungsgemäßes Verfahren zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Test-Kits zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Biochips bestimmt,
- einen Wirkstoff zur Aufrechterhaltung oder F\u00förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zust\u00e4nde der Haut einmal oder mehrmals auf die Haut aufbringt,
- c) erneut den Hautstatus durch ein erfindungsgemäßes Verfahren zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Test-Kits zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Biochips bestimmt, und
- d) die Wirksamkeit des Wirkstoffs durch den Vergleich der Ergebnisse aus
   a) und c) bestimmt.

Zur Beschleunigung des Testverfahrens ist es auch möglich, verschiedene Wirkstoffe oder Placebos parallel auf verschiedene Hautareale aufzubringen; beispielsweise einen Wirkstoff auf den linken Unterarm und ein Placebo auf den rechten Unterarm, oder umgekehrt.

Ein weiterer Gegenstand der vorliegenden Erfindung ist ein Test-Kit zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut in vitro, umfassend Mittel zur Durchführung des erfindungsgemäßen Testverfahrens.

Ein weiterer Gegenstand der vorliegenden Erfindung ist die Verwendung der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden zum Nachweis der Wirksamkeit von

kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom.

Ein weiterer Gegenstand der vorliegenden Erfindung ist ein Screening-Verfahren zur Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom in vitro, dadurch gekennzeichnet, daß man

- a) den Hautstatus durch ein erfindungsgemäßes Verfahren zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Test-Kits zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Biochips bestimmt.
- b) einen potentiellen Wirkstoff zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut einmal oder mehrmals auf die Haut aufbringt.
- c) erneut den Hautstatus durch ein erfindungsgemäßes Verfahren zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Test-Kits zur Bestimmung der Homeostase der Haut, oder mittels eines erfindungsgemäßen Biochips bestimmt, und
- d) wirksame Wirkstoffe durch den Vergleich der Ergebnisse aus a) und c) bestimmt.

Ein weiterer Gegenstand der vorliegenden Erfindung ist die Verwendung der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen. die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden, zur Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom.

Ein weiterer Gegenstand der vorliegenden Erfindung ist ein Verfahren zur Herstellung einer kosmetischen oder pharmazeutischen Zubereitung zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom, dadurch gekennzeichnet, daß man

- a) wirksame Wirkstoffe mit Hilfe des erfindungsgemäßen Screening-Verfahrens, oder der Verwendung zur Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut bestimmt und
- b) als wirksam befundene Wirkstoffe mit kosmetisch und pharmakologisch geeigneten und verträglichen Trägern vermischt.

# Tabellen:

Tabelle 6:

Nr.	Tag_Sequence	rel. Expr.	signific.	UniGene	Beschreibung
		In Haut		AccNr.	
1	ATCCGCGAGGC	45,00	63,99	Hs.180142	CLSP Calmodulin-like skin protei
2	GAGATAAATGA	23,00	31,88	Hs.3185	lymphocyte antigen 6 complex, locus D
3	TAAACCTGCTG	110,00	150,08	Hs.99923	lectin, galactoside-binding, soluble, 7 (galectin 7)
4	GATGTGCACGA	216,00	289,47	Hs.117729	keratin 14 (epidermolysis bullosa simplex, Dowling-M
5	ACATTTCAAAG	161,00	248,57	Hs.80828	keratin 1 (epidermolytic hyperkeratosis)
6	TTTGTAGAGGA	37,00			katanin p60 (ATPase-containing) subunit A 1
7	ACCTCCACTGG	139,00	214,56	Hs.112457	ESTs
	AATCTTGTTTC				ESTs
9	GAAAACAAAGT	467,00	100,00	Hs.99936	keratin 10 (epidermolytic hyperkeratosis; keratosis
10	GCCCCTGCTGA	148,00	181,69	Hs.195850	keratin 5 (epidermolysis bullosa simplex, Dowling-Me
11	CACACGGGCGA	26,00	32,16	Hs.194679	WNT1 inducible signaling pathway protein 2
12	GACAATAAATG	9,00	10,38	Hs.137556	Homo sapiens mRNA; cDNA DKFZp434A132 (from clone DKF
13	ACTACCATAAC	7,00	8,06	Hs.57929	slit (Drosophila) homolog 3
14	CATTGTAAATA	7,00	8,06	Hs.55279	protease inhibitor 5 (maspin)
15	ACCGGCGCCCG	29,00	32,51	Hs.65424	tetranectin (plasminogen-binding protein)
16	CCACCACGCTT	16,00	17,84		ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
17	CCCCGGCCACC	42,00	44,79	Hs.279604	(Manual assignment) desmin, muscle intermediate fila
18	TGAAATAAAAG	12,00	12,90		ESTs
19	ACTGAGTAGGT	4,00	4,39	Hs.38095	ATP-binding cassette, sub-family A (ABC1), member 8
20	ATCCTTGCTGA	26,00	26,12	Hs.2621	cystatin A (stefin A)
	CAGCCTGGGTG	5,00		Hs.171941	
	GATATGTTATA	5,00	5,10	Hs.117938	collagen, type XVII, alpha 1
23	TGGCTTCATCA	4,00	4,10	Hs.646	carboxypeptidase A3 (mast cell)
	CCTGTAACACC	3,00	3,09	Hs.74304	periplakin
	CCCCGGAGGTC	3,00			coagulation factor X
	AGATCAGTTGA	3,00		Hs.191805	
	CCCTCAGCACC	9,00			annexin A8
	CTTTATTCCAG	49,00			collagen, type I, alpha 1
	TCCACTGGCCT	23,00		Hs.57548	
30	CACGCAGTGGC	5,00	4,85	Hs.245545	EST

	TAGATTATATA	0.001	0.05	11. 400000	der til o
	TACATTATATA	3,00		Hs.198862	
32	ATGGATACGGG	10,00	9,05	Hs.250722	(Manual assignment) unclear, probably
<del></del>	0000000000			11 17000	reverse tag o
	CCGGGGGAGCC	43,00			collagen, type I, alpha 1
	CAGTTTTTTC	2,00			ESTs
	GTGGATTCAAG	2,00			NADPH oxidase 4
	TGTCTGTGTGT	2,00			ESTs
	TCTACACGTGC	2,00			properdin P factor, complement
	GAAATGGCAGT	2,00			ESTs
39	ACGAAACCTCG	2,00	1,84	Hs.285785	Homo sapiens cDNA FLJ20115 fis, clone
					COL05594
	GGCAATGCAGT	2,00		Hs.275505	
	CCTTTTCAGCA	2,00		Hs.25930	
	CCTCTTTAACA	2,00			ESTs
	TATCTAGCTGC	2,00	1,84	Hs.241545	hypothetical protein
44	GCTGTAATCCT	2,00	1,84	Hs.241382	tumor necrosis factor (ligand)
			<del></del>	<u> </u>	superfamily, member 1
45	GGGCAGCCGCC	2,00	1,84		CREB binding protein (Rubinstein-Taybi
<u></u>				2-2	syndrome)
46	CGCTTGTTTAA	2,00	1,84	Hs.180398	LIM domain-containing preferred
ļ	-				translocation partne
47	GCACACACCTG	2,00	1,84	Hs.171143	EST, Weakly similar to ALU1_HUMAN
40	CTCCACTCCCC	2 00	4.04	LI- 400074	ALU SUBFAMILY J SE
	CTGGAGTCGGC	2,00			Interleukin-1 Superfamily z
	GAGGTCAGTTG	2,00			DKFZP727G051 protein
	CCAGGCAAGAC GAAATCAAAAA	2,00			distal-less homeo box 3
	AATCTAGTTCT	13,00	22.70	Hs. 11/005	sialic acid binding Ig-like lectin 5
52	AATCIAGITCI	22,00	33,72		Human profilaggrin gene exons 1-3, 5' end
53	AAGCTAATAAA	9,00	7,89	Hs.88474	prostaglandin-endoperoxide synthase 1
					(prostaglandin
	TGTGCGGCTTC	5,00	4,43	Hs.162196	hypothetical protein FLJ20321
55	CAGGTTTCATA	66,00	54,72	Hs.24395	small inducible cytokine subfamily B
					(Cys-X-Cys), me
	CTGTCGTCATC	7,00	5,98	Hs.183860	hypothetical protein FLJ20277
57	ATAGCACGTGC	3,00	2,66		ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC
58	GTGAGAACTCG	3,00	2.66	Hs.250639	
	ACTTATTATGC	29,00			decorin
	CTTGCAGTCCT	5,00			Ris
	CATCTGTACTC	14,00			major histocompatibility complex, class
		, , , ,	,		II, DR beta
62	GTGGAGGGCAC	18,00	14,16		cystatin E/M
	AGGCAGGAAAA	5,00			ESTs, Weakly similar to hypothetical
					protein [H.sapi
64	AATTGAAAAGG	10,00	7,94	Hs.78344	myosin, heavy polypeptide 11, smooth
					muscle
	CTTTAAAATGA	3,00	2,50	Hs.8217	stromal antigen 2
66	TGTGCCAGTTT	3,00			ESTs, Weakly similar to ALU8_HUMAN
		1,5 -7			

					La caracteria de la car
				<del></del>	ALU SUBFAMILY SX
67	AGTAGCTGGGA	3,00	2,50	Hs.224534	EST, Weakly similar to alternatively spliced product
68	CCTCTGTCTCC	3,00	2,50	Hs.161031	Homo sapiens mRNA; cDNA DKFZp434K0322 (from clone DK
69	AACATTTAGGA	3,00	2.50	He 138380	KIAA0624 protein
	CAATAAAATTT	3,00			tumor protein 63 kDa with strong
					homology to p53
	GCCGCTCAAGG	3,00		Hs.126064	
	CCTGGTCAAGA	2,00			silver (mouse homolog) like
73	CCACCGCAGGA	2,00	1,68	Hs.85112	insulin-like growth factor 1 (somatomedia C)
74	ACACTTCTCAA	2,00	1,68	Hs.75652	glutathione S-transferase M5
75	CCTCTCTGGTC	2,00	1,68		heat shock 27kD protein family, member
					7 (cardiovasc
	GCATATCTGTG	2,00			KIAA1436 protein
77	AGCTGTGATGG	2,00		Hs.249983	
78	GCTAACTTAAA	2,00	1,68	Hs.20787	ESTs
79	CCTTGAAATCA	2,00	1,68	Hs.183161	ESTs
80	CTTTATCAATA	2,00	1,68	Hs.166017	microphthalmia-associated transcription
					factor
81	ACAGCCCTGAT	2,00			ribosomal protein L18a
82	GATACTCAGAA	2,00	1,68	Hs.144726	ESTs
83	GCCTGGGAGAC	2,00		Hs.118346	
84	TGGGTGGT	13,00	10,06		fragile X mental retardation, autosomal homolog 1
85	AGCTACCACAG	9,00	6,87		tenascin XB
	ACAGCGGCAAT	69,00			desmoplakin (DPI, DPII)
87	GTAAAATCCCA	3,00		Hs.278623	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
88	CACTTGTAATC	3,00	2.36		KIAA1185 protein
_	CTTGTAGTTCC	3,00			KIAA0677 gene product
90	GGGTTTTCTGG	3,00		Hs.153703	ESTs, Moderately similar to DHSA_HUMAN SUCCINATE DEH
91	CAGCAGAACTG	3,00	2,36		CGI-43 protein
92	CCACAGGAGAA	81,00		Hs.169902	solute carrier family 2 (facilitated glucose transpo
93	ATAGCCAGGGA	4,00	3.02		SRY (sex determining region Y)-box 20
	GTACAAAAGTA	4,00		Hs.9552	binder of Arl Two
-	TCACAGGGTCC	4,00			lamin A/C
	TTCTGTGTGCC	4,00			ESTs
	TAGCCGGGACG	10,00			Kruppel-like factor 2 (lung)
	ATCACACAGCT	6,00			leiomodin 1 (smooth muscle)
	ATCTCGAAAGG	8,00			hypothetical protein FLJ20037
	GACCCAACTGG	2,00			CD79B antigen (immunoglobulin-
		_,-,-5	-,-,		associated beta)
101	CACAGGGAGGA	2,00	1.54	Hs.84753	KIAA0246 protein
	CAGCTGGCCCA	2,00			fibulin 1
	GAGGGCTTTGC	2,00			aldo-keto reductase family 1, member C3

				,	
					(3-alpha hyd
104	GTGAGCCAAGA	2,00	1,54	Hs.75410	heat shock 70kD protein 5 (glucose- regulated protein
105	TACCCCAAAAA	2,00	1,54	Hs.6449	hypothetical protein FLJ20542
106	CAGGATGCTTG	2,00		Hs.56729	lymphocyte-specific protein 1
	AGTGTGTTGCA	2,00		Hs.56105	ESTs, Weakly similar to WDNM_RAT WDNM1 PROTEIN PRECU
108	AGGACCTGAAG	2,00	1,54	Hs.32352	hypothetical protein DKFZp434K1210
	ATTCTGGTCAT	2,00			ESTs, Weakly similar to dJ876B10.4 [H.sapiens]
110	CCTTTTGGGAG	2,00	1,54	Hs.186600	ESTs
111	CGGTTCATACA	2,00	1,54	Hs.169487	Kreisler (mouse) maf-related leucine zipper homolog
112	ATGGTGCCACC	2,00	1,54	Hs.161554	hypothetical protein FLJ20159
113	TAATGTTAATG	2,00			death-associated protein kinase 1
114	AGGGCCCTCTG	2,00	1,54	Hs.129014	hypothetical protein FLJ20207
115	GTGGCACGCGT	2,00	1,54	Hs.118243	deoxyribonuclease II, lysosomal
116	AGCTTGAGTTC	2,00	1,54	Hs.117582	CGI-43 protein
117	GTGGGGCCAAG	7,00			folate receptor 2 (fetal)
	GTGAAGCCTCA	5,00		Hs.271823	ESTs
	<u>ACCAGACAGAC</u>	3,00			ESTs
120	GTGAAACTCTT	3,00	2,23		RNA POLYMERASE I AND
					TRANSCRIPT RELEASE FACTOR
	ATTTCCATTAA	3,00	2,23	Hs.284126	hairless (mouse) homolog
:	GTGGTAAGCAC	3,00			ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
	GTTTTGCCCAC	3,00	2,23	Hs.151407	cartilage intermediate layer protein, nucleotide pyr
124	GCCCACACAGC	8,00	5,71	Hs.1690	heparin-binding growth factor binding protein
125	TTTCCTCTCAA	38,00	25,30	Hs.184510	stratifin
126	CGGGAGCGCTA	13,00	8,84		ESTs, Weakly similar to AF208846_1 BM-004 [H.sapiens
	TTGCATATCAG	10,00		Hs.82237	ataxia-telangiectasia group D-associated protein
128	AGGCCTCGGCA	5,00	3,55	Hs.286202	Homo sapiens cDNA FLJ11346 fis, clone PLACE1010900
	GTGGCGAATGA	12,00			desmocollin 1
	TGTGAAGCCTT	12,00		Hs.5476	serine protease inhibitor, Kazal type, 5
	TCAGACTTTTG	9,00		Hs.5889	ESTs, Weakly similar to AC004876_5 similar to predic
132	ATTTCTTCAAG	9,00		Hs.31386	ESTs, Highly similar to JE0174 frizzled protein-2 -
	GAATTATACTT	10,00	6,73	Hs.104800	hypothetical protein FLJ10134
	TCTGGGGAACA	3,00	2,12	Hs.184390	similar to aspartate beta hydroxylase (ASPH)
	GCAAAAACCCG	3,00			ribosomal protein L37a
136	AATGTTGTGCA	4,00	2,78	Hs.91546	cytochrome P450 retinoid metabolizing protein

	ACAATGTTGTA	2,00		Hs.7678	cellular retinoic acid-binding protein 1
	CGAGAGTGTGA	2,00			ESTs
	GTATAAAAAAA	2,00			hypothetical protein FLJ20623
	AGGTCGAGGCT	2,00		Hs.270125	
141	CCCGGCCCAGT	2,00	1,43	Hs.243324	EST, Weakly similar to ALUA_HUMAN
					IIII ALU CLASS A W
	TTGACCCAGCC	2,00		Hs.193745	
	TATTTTATTTG	2,00			purinergic receptor (family A group 5)
144	GCATCATAGGT	2,00	1,43	Hs.184108	ribosomal protein L21 (gene or
					pseudogene)
145	TACCGCTCCCT	2,00	1,43		Homo sapiens mRNA; cDNA
					DKFZp434G2416 (from clone DK
146	CTCCTGTGGTC	2,00	1,43	Hs.169851	ESTs, Weakly similar to ALU4_HUMAN
L					ALU SUBFAMILY SB2
147	GGTGTCTCCTC	2,00		Hs.146038	
148	CAATCTTGTGA	2,00		Hs.104353	
149	CCTACAAAAA	1,00			ESTs
150	TGTGCCCAGCC	1,00	0,69	Hs.97905	ovo (Drosophila) homolog-like 1
151	GATGGGGACAG	1,00	0,69	Hs.92195	ESTs
152	TAAAAATATTG	1,00	0,69	Hs.89695	insulin receptor
153	TTTAAGTTAGG	1,00	0,69	Hs.82932	cyclin D1 (PRAD1: parathyroid
					adenomatosis 1)
154	GTATATGTATT	1,00	0,69	Hs.7917	DKFZP564K247 protein
155	CTAAAGTGTCA	1,00	0,69	Hs.7910	RING1 and YY1 binding protein
156	AATTTGGCTTT	1,00	0,69	Hs.7734	Homo sapiens cDNA FLJ20684 fis, clone
					KAIA3469
157	CCGTGGCACCA	1,00	0,69	Hs.77208	ESTs
158	AGCACCCTTGT	1,00	0,69	Hs.75871	protein kinase C binding protein 1
159	CCTTTGAGAGC	1,00	0,69		hypothetical protein
160	TAAATGTAAAT	1,00	0,69		KIAA0427 gene product
161	TACAGACATAC	1,00	0,69	Hs.63984	cadherin 13, H-cadherin (heart)
162	GTAGCATTTGC	1,00	0,69	Hs.63302	myotubularin related protein 3
163	AAGTAGGTTTT	1,00	0,69	Hs.50216	zinc finger protein (ZFD25)
164	ATCACTCCCCA	1,00	0,69	Hs.37058	calcitonin/calcitonin-related polypeptide,
					alpha
165	TAGGCAGACCT	1,00	0,69	Hs.35488	ESTs, Moderately similar to
					ALU6_HUMAN ALU SUBFAMILY
	GCCTGGCCAGG	1,00		Hs.3343	phosphoglycerate dehydrogenase
167	GTCATCTTGTT	1,00	0,69	Hs.32366	ESTs, Moderately similar to
					TWST_HUMAN TWIST RELATED
168	CTCAACAACCA	1,00	0,69	Hs.30036	ESTs, Weakly similar to ALU1_HUMAN
					ALU SUBFAMILY J S
	TTATATTAATA	1,00		Hs.29205	alpha integrin binding protein 63
170	GATTAAACCTT	1,00	0,69	Hs.287357	ESTs, Moderately similar to meningioma-
					expressed ant
171	GATGGAGGTTA	1,00	0,69	Hs.285224	ESTs, Weakly similar to unnamed
L				_	protein product [H.s
	TACAGGCGTGG	1,00		Hs.283329	
I 173	TGTGTGTGTAT	1,00	0,69	Hs.278676	Homo sapiens mRNA; cDNA

174   GGTGCCTGTAA						
IIII ALU CLASS C   1,00    0,69   Hs.278242   tubulin, alpha, ubiquitous   176   ACAGGAGCGTT    1,00    0,69   Hs.275996   EST   177   ATGCCTACTCT    1,00    0,69   Hs.274934   EST   178   AGCACAATCTT    1,00    0,69   Hs.274934   EST   178   AGCACAATCTT    1,00    0,69   Hs.274128   Homo sapiens cDNA FLJ10131 fis, clone   HEMBA1003041   179   GTTCTGTTTGG    1,00    0,69   Hs.271040   ESTs   180   TTGTGATTATA    1,00    0,69   Hs.250320   TRABID protein   181   TATGCAGATCA    1,00    0,69   Hs.250320   TRABID protein   181   TATGCAGATCA    1,00    0,69   Hs.250520   ESTs   182   GTTGAGGACAT    1,00    0,69   Hs.250520   ESTs   183   GCCAGTGGCTG    1,00    0,69   Hs.242481   ESTs   183   GCCAGTGGCTG    1,00    0,69   Hs.242481   ESTs   185   CAAAAAGTTGA    1,00    0,69   Hs.242481   ESTs   185   CAAAAAGTTGA    1,00    0,69   Hs.23303   Homo sapiens mRNA; cDNA DKFZp434M2217 (from clone DK    186   GCCACAGCTGG    1,00    0,69   Hs.233013   EST   190   CAAAAGAATAA    1,00    0,69   Hs.233193   EST   191   TAACAGTAATA    1,00    0,69   Hs.233193   EST   191   TAACAGATAATA    1,00    0,69   Hs.233193   EST   192   TGCCTAGGAAA    1,00    0,69   Hs.226356   DKFZp586F1922 (from clone DK    193   CCCTAGGAGAC    1,00    0,69   Hs.211258   ESTs   195   CATTCTCTT    1,00    0,69   Hs.203750   EST   195   CATTCTCTT    1,00    0,69   Hs.207181   ESTS   198   CATTCTCTTT    1,00    0,69   Hs.207181   ESTS   199   CCTGGAGTGCA    1,00    0,69   Hs.207181   ESTS   199   CATTCTCAGCCT    1,00    0,69   Hs.207180   EST   199   CTGGAGTGCA    1,00    0,69   Hs.17630   EST   199   CTGGGAGTGCA    1,00    0,69   Hs.17630   EST   190   CTGGGATGCA    1,00    0,69   Hs.17630						DKFZp434J1630 (from clone DK
175  AAGTTGTGGCC	174	GGTGCCTGTAA	1,00	0,69		
176   ACAGGAGCGTT	175	AAGTTGTGGCC	1.00	0.69		
1778   AGCACACTCT						
178   AGCACAATCTT						
HEMBA1003041   HEMB						
180   TTGTGATTATA						HEMBA1003041
181   TATGCAGATCA						
182 GTTGAGGACAT						
183   GCCAGTGGCTG						
184   GCAGATCTTCC						
185   TIGAAACITICT						
186   CAAAAAGTTGA						
188   GCCACAGCTGG	-					
DKFZp434M2217 (from clone DK	-					
188   GCCACAGCTGG	187	CTCCATTCTCA	1,00	0,69		
189   GGGAAAGAAGG						
190   CAAAAGAATAA	188	GCCACAGCTGG	1,00	0,69	Hs.23565	ESTs
191   TAACAGTAATA   1,00   0,69   Hs.231913   ESTs     192   TGCCTAGGAAA   1,00   0,69   Hs.226356   Homo sapiens mRNA; cDNA DKFZp586F1922 (from clone DK     193   CCCTAGGAGAC   1,00   0,69   Hs.217484   ESTs     194   GGGTTGTTGTA   1,00   0,69   Hs.2017181   ESTs     195   AACCCCCAAAC   1,00   0,69   Hs.2017181   ESTs     196   CTTTTTCTTT   1,00   0,69   Hs.203750   EST     197   TGGGAATTGTG   1,00   0,69   Hs.201306   ESTs     198   CATTCCAGCCT   1,00   0,69   Hs.201306   ESTs     199   GCTGGAGTGCA   1,00   0,69   Hs.195484   Homo sapiens mRNA full length insert cDNA clone EURO     200   GAAGGCCAGCT   1,00   0,69   Hs.193488   Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK     201   ACGTATTTGAG   1,00   0,69   Hs.18800   hypothetical protein FLJ20281     203   CTAAATGTGAA   1,00   0,69   Hs.181163   high-mobility group (nonhistone chromosomal) protein     204   CCTGGCTCTAA   1,00   0,69   Hs.178705   Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616     205   CTGTCCTTGTT   1,00   0,69   Hs.176333   ESTs     206   CAATATTTGAG   1,00   0,69   Hs.170673   ESTs, Weakly similar to AF126780_1 retinal short-cha     208   GTCCTTGACCA   1,00   0,69   Hs.170673   ESTs, Weakly similar to AF126780_1 retinal short-cha     209   GCAATGACCTG   1,00   0,69   Hs.170162   KIAA1357 protein	189	GGGAAAGAAGG	1,00	0,69	Hs.233193	EST
192   TGCCTAGGAAA	190	CAAAAGAATAA	1,00	0,69	Hs.233013	EST
DKFZp586F1922 (from clone DK	191	TAACAGTAATA	1,00			
193 CCCTAGGAGAC         1,00         0,69 Hs.217484 ESTs           194 GGGTTGTTGTA         1,00         0,69 Hs.211258 ESTs           195 AACCCCCAAAC         1,00         0,69 Hs.207181 ESTs           196 CTTTTTCTTT         1,00         0,69 Hs.204917 EST           197 TGGGAATTGTG         1,00         0,69 Hs.201306 ESTs           198 CATTCCAGCCT         1,00         0,69 Hs.201306 ESTs           199 GCTGGAGTGCA         1,00         0,69 Hs.195484 Homo sapiens mRNA full length insert cDNA clone EURO           200 GAAGGCCAGCT         1,00         0,69 Hs.194624 ESTs           201 ACGTATTTGAG         1,00         0,69 Hs.193488 Homo sapiens mRNA; cDNA DKFZP434O1521 (from clone DK           202 TAATTTAAACC         1,00         0,69 Hs.18800 hypothetical protein FLJ20281           203 CTAAATGTGAA         1,00         0,69 Hs.181163 high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.176333 ESTs           205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           207 CAATGGATGGC         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG	192	TGCCTAGGAAA	1,00	0,69	Hs.226356	Homo sapiens mRNA; cDNA
194 GGGTTGTTGTA         1,00         0,69 Hs.211258 ESTs           195 AACCCCCAAAC         1,00         0,69 Hs.207181 ESTs           196 CTTTTTTCTTT         1,00         0,69 Hs.204917 EST           197 TGGGAATTGTG         1,00         0,69 Hs.203750 EST           198 CATTCCAGCCT         1,00         0,69 Hs.201306 ESTs           199 GCTGGAGTGCA         1,00         0,69 Hs.195484 Homo sapiens mRNA full length insert cDNA clone EURO           200 GAAGGCCAGCT         1,00         0,69 Hs.193488 Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK           201 ACGTATTTGAG         1,00         0,69 Hs.18800 hypothetical protein FLJ20281           202 TAATTTAAACC         1,00         0,69 Hs.181163 high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.178705 Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.174030 a disintegrin and metalloproteinase domain 28           207 CAATGGATGGC         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.17052 interleukin 9 receptor           210 TCCTAATTCAG         1,00 <td></td> <td></td> <td></td> <td></td> <td></td> <td>DKFZp586F1922 (from clone DK</td>						DKFZp586F1922 (from clone DK
195   AACCCCCAAAC   1,00   0,69   Hs.207181   ESTs     196   CTTTTTCTTT   1,00   0,69   Hs.204917   EST     197   TGGGAATTGTG   1,00   0,69   Hs.203750   EST     198   CATTCCAGCCT   1,00   0,69   Hs.201306   ESTs     199   GCTGGAGTGCA   1,00   0,69   Hs.195484   Homo sapiens mRNA full length insert cDNA clone EURO     200   GAAGGCCAGCT   1,00   0,69   Hs.194624   ESTs     201   ACGTATTTGAG   1,00   0,69   Hs.193488   Homo sapiens mRNA; cDNA DKFZP434O1521 (from clone DK     202   TAATTTAAACC   1,00   0,69   Hs.18800   hypothetical protein FLJ20281     203   CTAAATGTGAA   1,00   0,69   Hs.181163   high-mobility group (nonhistone chromosomal) protein     204   CCTGGCTCTAA   1,00   0,69   Hs.178705   Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616     205   CTGTCCTTGTT   1,00   0,69   Hs.176333   ESTs     206   CAATATTTGAG   1,00   0,69   Hs.170673   ESTs, Weakly similar to AF126780_1 retinal short-cha     208   GTCCTTGACCA   1,00   0,69   Hs.170524   ESTs     209   GCAATGACCTG   1,00   0,69   Hs.170162   KIAA1357 protein						
196   CTTTTTCTTT   1,00   0,69   Hs.204917   EST     197   TGGGAATTGTG   1,00   0,69   Hs.203750   EST     198   CATTCCAGCCT   1,00   0,69   Hs.201306   ESTs     199   GCTGGAGTGCA   1,00   0,69   Hs.195484   Homo sapiens mRNA full length insert cDNA clone EURO     200   GAAGGCCAGCT   1,00   0,69   Hs.194624   ESTs     201   ACGTATTTGAG   1,00   0,69   Hs.193488   Homo sapiens mRNA; cDNA DKFZP434O1521 (from clone DK     202   TAATTTAAACC   1,00   0,69   Hs.18800   hypothetical protein FLJ20281     203   CTAAATGTGAA   1,00   0,69   Hs.181163   high-mobility group (nonhistone chromosomal) protein     204   CCTGGCTCTAA   1,00   0,69   Hs.178705   Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616     205   CTGTCCTTGTT   1,00   0,69   Hs.176333   ESTs     206   CAATATTTGAG   1,00   0,69   Hs.174030   a disintegrin and metalloproteinase domain 28     207   CAATGGATGGC   1,00   0,69   Hs.170673   ESTs, Weakly similar to AF126780_1 retinal short-cha     208   GTCCTTGACCA   1,00   0,69   Hs.170524   ESTs     209   GCAATGACCTG   1,00   0,69   Hs.170162   KIAA1357 protein	194	GGGTTGTTGTA	1,00	0,69	Hs.211258	ESTs
197 TGGGAATTGTG         1,00         0,69 Hs.203750 EST           198 CATTCCAGCCT         1,00         0,69 Hs.201306 ESTs           199 GCTGGAGTGCA         1,00         0,69 Hs.195484 Homo sapiens mRNA full length insert cDNA clone EURO           200 GAAGGCCAGCT         1,00         0,69 Hs.194624 ESTs           201 ACGTATTTGAG         1,00         0,69 Hs.193488 Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK DKFZp434O1521 (from clone DK PLZ0281 Ingh-mobility group (nonhistone chromosomal) protein           202 TAATTTAAACC         1,00         0,69 Hs.181163 high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.178705 Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.174030 a disintegrin and metalloproteinase domain 28           207 CAATGGATGGC         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.170162 KIAA1357 protein	195	AACCCCCAAAC	1,00	0,69	Hs.207181	ESTs
198 CATTCCAGCCT         1,00         0,69 Hs.201306 ESTs           199 GCTGGAGTGCA         1,00         0,69 Hs.195484 Homo sapiens mRNA full length insert cDNA clone EURO           200 GAAGGCCAGCT         1,00         0,69 Hs.194624 ESTs           201 ACGTATTTGAG         1,00         0,69 Hs.193488 Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK DKFZp434O1521 (from clone DK DKFZp434O1521 (from clone DK DKFZp434O1521)           202 TAAATTTAAACC         1,00         0,69 Hs.18800 hypothetical protein FLJ20281           203 CTAAATGTGAA         1,00         0,69 Hs.181163 high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.178705 Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.1702 interleukin 9 receptor           210 TCCTAATTCAG         1,00         0,69 Hs.170162 KIAA1357 protein				0,69	Hs.204917	EST
199         GCTGGAGTGCA         1,00         0,69         Hs.195484         Homo sapiens mRNA full length insert cDNA clone EURO           200         GAAGGCCAGCT         1,00         0,69         Hs.194624         ESTs           201         ACGTATTTGAG         1,00         0,69         Hs.193488         Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK           202         TAATTTAAACC         1,00         0,69         Hs.18800 hypothetical protein FLJ20281           203         CTAAATGTGAA         1,00         0,69         Hs.181163 high-mobility group (nonhistone chromosomal) protein           204         CCTGGCTCTAA         1,00         0,69         Hs.178705 Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205         CTGTCCTTGTT         1,00         0,69         Hs.176333         ESTs           206         CAATATTTGAG         1,00         0,69         Hs.174030 a disintegrin and metalloproteinase domain 28           207         CAATGGATGGC         1,00         0,69         Hs.170673         ESTs, Weakly similar to AF126780_1 retinal short-cha           208         GTCCTTGACCA         1,00         0,69         Hs.170524         ESTs           209         GCAATGACCTG         1,00         0,69         Hs.170162         KIAA1357 protein	197	TGGGAATTGTG	1,00	0,69	Hs.203750	EST
CDNA clone EURO	198	CATTCCAGCCT	1,00			
200 GAAGGCCAGCT         1,00         0,69 Hs.194624 ESTs           201 ACGTATTTGAG         1,00         0,69 Hs.193488 Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK           202 TAATTTAAACC         1,00         0,69 Hs.18800 hypothetical protein FLJ20281           203 CTAAATGTGAA         1,00         0,69 Hs.181163 high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.178705 Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.174030 a disintegrin and metalloproteinase domain 28           207 CAATGGATGGC         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.1702 interleukin 9 receptor           210 TCCTAATTCAG         1,00         0,69 Hs.170162 KIAA1357 protein	199	GCTGGAGTGCA	1,00	0,69	Hs.195484	
201 ACGTATTTGAG         1,00         0,69 Hs.193488 Homo sapiens mRNA; cDNA DKFZp434O1521 (from clone DK           202 TAATTTAAACC         1,00         0,69 Hs.18800 hypothetical protein FLJ20281           203 CTAAATGTGAA         1,00         0,69 Hs.181163 high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.178705 Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.174030 a disintegrin and metalloproteinase domain 28           207 CAATGGATGGC         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.1702 interleukin 9 receptor           210 TCCTAATTCAG         1,00         0,69 Hs.170162 KIAA1357 protein	200	GAAGGCCAGCT	1,00	0,69	Hs.194624	
DKFZp434O1521 (from clone DK						
202 TAATTTAAACC         1,00         0,69 Hs.18800         hypothetical protein FLJ20281           203 CTAAATGTGAA         1,00         0,69 Hs.181163         high-mobility group (nonhistone chromosomal) protein           204 CCTGGCTCTAA         1,00         0,69 Hs.178705         Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205 CTGTCCTTGTT         1,00         0,69 Hs.176333         ESTs           206 CAATATTTGAG         1,00         0,69 Hs.174030         a disintegrin and metalloproteinase domain 28           207 CAATGGATGGC         1,00         0,69 Hs.170673         ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524         ESTs           209 GCAATGACCTG         1,00         0,69 Hs.170162         KIAA1357 protein						
203         CTAAATGTGAA         1,00         0,69         Hs.181163         high-mobility group (nonhistone chromosomal) protein           204         CCTGGCTCTAA         1,00         0,69         Hs.178705         Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205         CTGTCCTTGTT         1,00         0,69         Hs.176333         ESTs           206         CAATATTTGAG         1,00         0,69         Hs.174030         a disintegrin and metalloproteinase domain 28           207         CAATGGATGGC         1,00         0,69         Hs.170673         ESTs, Weakly similar to AF126780_1 retinal short-cha           208         GTCCTTGACCA         1,00         0,69         Hs.170524         ESTs           209         GCAATGACCTG         1,00         0,69         Hs.170162         KIAA1357 protein	202	TAATTTAAACC	. 1,00	0,69	Hs.18800	
Chromosomal) protein	203	CTAAATGTGAA		0,69	Hs.181163	high-mobility group (nonhistone
204         CCTGGCTCTAA         1,00         0,69         Hs.178705         Homo sapiens cDNA FLJ11333 fis, clone PLACE1010616           205         CTGTCCTTGTT         1,00         0,69         Hs.176333         ESTs           206         CAATATTTGAG         1,00         0,69         Hs.174030         a disintegrin and metalloproteinase domain 28           207         CAATGGATGGC         1,00         0,69         Hs.170673         ESTs, Weakly similar to AF126780_1 retinal short-cha           208         GTCCTTGACCA         1,00         0,69         Hs.170524         ESTs           209         GCAATGACCTG         1,00         0,69         Hs.1702         interleukin 9 receptor           210         TCCTAATTCAG         1,00         0,69         Hs.170162         KIAA1357 protein				·		
PLACE1010616	204	CCTGGCTCTAA	1,00	0,69	Hs.178705	
205 CTGTCCTTGTT         1,00         0,69 Hs.176333 ESTs           206 CAATATTTGAG         1,00         0,69 Hs.174030 a disintegrin and metalloproteinase domain 28           207 CAATGGATGGC         1,00         0,69 Hs.170673 ESTs, Weakly similar to AF126780_1 retinal short-cha           208 GTCCTTGACCA         1,00         0,69 Hs.170524 ESTs           209 GCAATGACCTG         1,00         0,69 Hs.1702 interleukin 9 receptor           210 TCCTAATTCAG         1,00         0,69 Hs.170162 KIAA1357 protein				·		
206         CAATATTTGAG         1,00         0,69         Hs.174030         a disintegrin and metalloproteinase domain 28           207         CAATGGATGGC         1,00         0,69         Hs.170673         ESTs, Weakly similar to AF126780_1 retinal short-cha           208         GTCCTTGACCA         1,00         0,69         Hs.170524         ESTs           209         GCAATGACCTG         1,00         0,69         Hs.1702         interleukin 9 receptor           210         TCCTAATTCAG         1,00         0,69         Hs.170162         KIAA1357 protein	205	CTGTCCTTGTT	1,00	0,69	Hs.176333	
domain 28						
retinal short-cha						
retinal short-cha	207	CAATGGATGGC	1,00	0,69	Hs.170673	
209 GCAATGACCTG         1,00         0,69 Hs.1702         interleukin 9 receptor           210 TCCTAATTCAG         1,00         0,69 Hs.170162         KIAA1357 protein						retinal short-cha
209 GCAATGACCTG         1,00         0,69 Hs.1702         interleukin 9 receptor           210 TCCTAATTCAG         1,00         0,69 Hs.170162         KIAA1357 protein	208	GTCCTTGACCA	1,00	0,69	Hs.170524	ESTs
210 TCCTAATTCAG 1,00 0,69 Hs.170162 KIAA1357 protein	209	GCAATGACCTG				
	210	TCCTAATTCAG	1,00	0,69		
	211	TTAATGATCTT	1,00			

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212	TTAAGTGTTCT	1,00	0.69	He 159239	toll-like receptor 4
	TGAAGCGTTTA	1,00			protein tyrosine phosphatase, non-
213	10,410001117	1,00	0,00	1 13. 100000	receptor type 21
214	GTGGAGCTTAA	1,00	0.60	Hs.152385	
	GAGACTGGGGC	1,00		Hs.142854	
<b></b>	GTGGTACTCGC	1,00			ESTs, Weakly similar to S59501
210	GIGGIACICGC	1,00	0,09		interferon receptor J
217	TGCAGTGCTTG	1,00	0.60		chloride channel 2
	AGCTTATTGGC	1,00			KIAA1098 protein
-	ACTICTGCTTA	1,00		Hs.13740	
	GATAGAAATTT	1,00		Hs.131987	the state of the s
	TTTGACTAATT	1,00		Hs.131761	
	TCAAAATGACA	1,00		Hs. 131272	
$\overline{}$	CCAGCTAGTTT			Hs. 128692	
		1,00			
	TAGAGGAGTTG	1,00		Hs. 125815	
	GGGAAACACCA	1,00		Hs.123471	
	GAAATGAGTGT	1,00		Hs.1200	arachidonate 12-lipoxygenase
	ATCTTGGTACT	1,00			fibronectin 1
	CATATCATCTC	1,00		Hs.118130	
	ATGTGACTTTT	1,00			CGI-43 protein
	CCACTGTAAGC	1,00	0,69	Hs.117582	CGI-43 protein
231	TATAAGGCTGA	1,00			ESTs, Weakly similar to ZN84_HUMAN ZINC FINGER PROTE
232	GCCACTGCCAC	1,00	0,69		S100 calcium-binding protein A9
$ldsymbol{ldsymbol{ldsymbol{eta}}}$					(calgranulin B)
	TGAGGCCAGGG	1,00	0,69	Hs.110128	hypothetical protein FLJ10060
	GTTTATTTGAA	1,00		Hs.109087	
	AAAATTGTTAG	1,00			hypothetical protein FLJ20129
	GTGATGGGCTC	9,00			envoplakin
237	AGGCTCCTGGC	52,00	31,57	Hs.24395	small inducible cytokine subfamily B (Cys-X-Cys), me
238	AACAGCAAGGA	5,00	3,33	Hs.20665	ESTs
239	CCACGGGATTC	14,00	8,79		collagen, type III, alpha 1 (Ehlers-Danios syndrome
240	GATTTCGTTTT	4,00	2,68	Hs.738	early growth response 1
241	TATAGCCCTCA	4,00	2,68	Hs.64311	a disintegrin and metalloproteinase domain 17 (tumor
242	GTGACACGTGC	4,00	2,68	Hs.282996	ESTs
243	AGCTGTCGTAG	3,00	2,02	Hs.86674	ESTs
244	ATTGTTTCAAG	3,00			ESTs, Moderately similar to TWST HUMAN TWIST RELATED
245	AGACCCTGTCT	3,00	2.02	Hs.239283	
	TTGGCAAGGCT	3,00		Hs.184720	
	ATCATAGCTCA	6,00		Hs.97876	ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
2/19	CCTACCACCAT	11,00	6 72	Hs.8468	RelA-associated inhibitor
	CCAGGGCAACA	40,00			(Manual assignment) ORF-less transcript
					in MEN1 regi
250	TACAGTATTTT	2,00	<u>1,3</u> 3	Hs.82921_	solute carrier family 35 (CMP-sialic acid

					transporte
251	TGCCAGGTGCA	2,00	1 33	Hs.75442	
	TITITIATTCC	2,00			interleukin 10 receptor, alpha
	AATATTTTAT	2,00			ESTs, Highly similar to JE0174 frizzled
					protein-2 -
254	GAGAACCACCT	2,00	1,33	Hs.30250	v-maf musculoaponeurotic fibrosarcoma (avian) oncoge
255	GACACACCGAA	2,00	1,33	Hs.274243	receptor tyrosine kinase-like orphan receptor 1
256	GGCTTGTCTAT	2,00	1,33		ESTs, Weakly similar to weak similarity to HSP90 [C.
257	GGCAATATAGT	2,00	1,33		ESTs, Weakly similar to unknown protein [H.sapiens]
258	AGGATAACTTC	2,00	1,33	Hs.184482	DKFZP586D0624 protein
259	GATCAATCAGT	2,00	•		small inducible cytokine subfamily A (Cys-Cys), memb
260	TTGAATATTAA	2,00	1,33	Hs.161554	hypothetical protein FLJ20159
261	TCCTCACTTCA	2,00	1,33	Hs.158455	ESTs
262	GTTGCAGCATT	2,00			HYA22 protein
263	GTGTCTGTCTC	2,00		Hs.137432	
	GGCCTCTCCGA	2,00			hematopoietic protein 1
	ACAGAATGCCT	17,00		Hs.79732	
	GGGGCTGCCCA	15,00			tumor endothelial marker 1 precursor
	AAAATCGCTTG	5,00			COX15 (yeast) homolog, cytochrome c
					oxidase assembly
_	GTATAAACGTC	3,00			stromal cell-derived factor 1
$\overline{}$	CACTTTACCAG	3,00			runt-related transcription factor 3
	TTAATTACAGT	3,00			serum/glucocorticoid regulated kinase
	TCAGCGACCCT	4,00	2,49	Hs.169946	GATA-binding protein 3
	ATGGCACATTC	4,00			Homo sapiens mRNA; cDNA DKFZp762O124 (from clone DKF
273	AGGCTCAGGTC	9,00	13,62	Hs.78344	myosin, heavy polypeptide 11, smooth muscle
274	AACAGGGGCCA	5,00	3,05	Hs.262958	ESTs, Weakly similar to alternatively spliced produc
275	CTGAAATCTAT	5,00	3,05	Hs.253467	
	GAGAAATCCCG	5,00		Hs.150298	
277	TCAAAAGACCT	12,00			v-fos FBJ murine osteosarcoma viral oncogene homolog
278	GAAAACAAACA	2,00	1.25	Hs.83004	interleukin 14
	TGCATCTGTAC	2,00		Hs.58589	glycogenin 2
	ATGAAACTCCA	2,00		Hs.277951	
	TITATTCCTCT	2,00			ESTs
	GGGAGACCTGT	2,00			EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SE
283	CACTATGTAAA	2,00	1,25	Hs.24143	Wiskott-Aldrich syndrome protein interacting protein
284	TACAGCGGCAG	2,00	1,25	Hs.205616	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S

005	A OTOGOTTOGO		4.05	11. 450004	I=0=
	AGTGCCTTGGG	2,00		Hs.178604	
	CGATGCTGACG	2,00			hypothetical protein FLJ20159
287	GTGGTGTAATC	3,00	1,84	Hs.261734	ESTs, Moderately similar to
<u></u>				ļ	ALU7_HUMAN ALU SUBFAMILY
	TATCCCAGAAT	3,00		Hs.175819	
289	TCTAAAAAGGC	3,00	1,84	Hs.16622	zinc finger protein 185 (LIM domain)
290	TCGAAACGCTG	3,00	1,84	Hs.136528	ESTs, Moderately similar to
					ALU1_HUMAN ALU SUBFAMILY
	ACATTCTTTTT	22,00	11,94	Hs.82226	glycoprotein (transmembrane) nmb
292	CTGTTTGTTCA	8,00	4,59	Hs.211582	myosin, light polypeptide kinase
293	CAGTACTGTAT	4,00	2,41	Hs.9295	elastin (supravalvular aortic stenosis,
					Williams-Beu
294	GGTGAAACCCC	4,00	2,41	Hs.284878	EST
	GCCAAACCCCA	4,00		Hs.194264	
	GCCGACGCCAG	4,00		Hs.165565	
	GAAGAGGACAA	5,00		Hs.120451	
	TAAATAAGAAA	1,00			zona pellucida binding protein
	AAAGATCCCTC	1,00			ESTs
	ATCAAAGGTTA	1,00			Homo sapiens mRNA; cDNA
555	, , , , , , , , , , , , , , , , , , , ,	.,,55	0,01	110.04700	DKFZp564O222 (from clone DKF
301	GCAAGGTTGGT	1,00	0.61	Hs.94761	ESTs, Weakly similar to KIAA0561
00.	00,110011001	.,00	0,0 .	113.04701	protein [H.sapiens]
302	GGGACAAAAA	1,00	0.61	Hs.93788	ESTs .
	ACTACCTCTGA	1,00			hypothetical protein FLJ20163
	GCTGGGCCCAG	1,00		Hs.90964	Homo sapiens cDNA FLJ20812 fis, clone
	00100000000	1,00	0,01		ADSE01316
305	CAAAGAAATAG	1,00	0.61		ESTs
	AATACAAGTAT	1,00			KIAA1301 protein
	CTCTGTGGCTC	1,00			insulin-like growth factor 1 (somatomedia
00,	01010100010	1,00	0,01	115.05112	C)
308	TATTTGAAAGT	1,00	0.61	Hs.82664	ETAA16 protein
	CCCGCCTCCGT	1,00			Cbp/p300-interacting transactivator, with
ا	00000010001	1,00	0,01	113.02071	Glu/Asp-ri
310	TCACCGTAGCC	1,00	0.61	Hs.82042	solute carrier family 23 (nucleobase
5,5	. 5. 1000 17 1000	.,55	0,01	113.02072	transporters),
311	CCAAAAATTAA	1,00	0.61	Hs.81424	ubiquitin-like 1 (sentrin)
	ACTCGTATATG	1,00			
	TCAAGATGAAG	1,00		Hs.78948	interleukin 1 receptor antagonist Rab geranylgeranyltransferase, beta
5,5		1,50	0,01	1 13.7 0340	subunit
314	CGTCTATCCAT	1,00	0.61	Hs.76084	lamin B2
$\rightarrow$	AAAGTGAAATG	1,00		Hs.75912	KIAA0257 protein
<del></del>	TTCTGGTGCTG	1,00			
	CTGGCGTCGTC				transgelin 2
-		1,00		Hs.75640	natriuretic peptide precursor A
$\overline{}$	GGCTGGGGAGG	1,00		Hs.75061	MARCKS-like protein
	TGCCCCCAAAA	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	CTITOCTCATT	1,00			cytochrome c oxidase subunit VIc
	CTTTCCTCATT	1,00		Hs.7381	voltage-dependent anion channel 3
	TTGGACTGAGG	1,00			ganglioside expression factor 2
323	AGCATTAAAAA	1,00	0,61	Hs.61638	myosin X

324	CTGCTTTAAAA	1,00			brain-derived neurotrophic factor
325	TGCTTAAAAAT	1,00		Hs.5534	ESTs
326	ATGAACCCCCT	1,00		Hs.5011	RNA binding motif protein 9
327	TGAAACTTCCC	1,00	0,61	Hs.4994	transducer of ERBB2, 2
328	CAAACAAAAA	1,00	0,61	Hs.43728	hypothetical protein
329	GAGGTAACTAC	1,00	0,61	Hs.43712	ESTs
330	CATTAAAAAAAT	1,00	0,61	Hs.36908	activating transcription factor 1
331	CAATGGTGAAA	1,00	0,61	Hs.35093	lymphoid blast crisis oncogene
332	CTAATGAATGT	1,00	0,61	Hs.29809	Homo sapiens mRNA; cDNA
					DKFZp434C185 (from clone DKF
333	AGGGCTTTCAC	1,00	0,61	Hs.29797	ribosomal protein L10
334	TAACCGTGGAA	1,00	0,61	Hs.29647	uncharacterized hematopoietic
1				·	stem/progenitor cells
335	TCTCCCACACC	1,00	0,61	Hs.2961	S100 calcium-binding protein A3
336	CCTGGATCTCC	1,00	0,61		ESTs
337	GGCCTGGCACT	1,00	0,61	Hs.283388	ESTs
338	ACTITGTTTTT	1,00	0,61	Hs.28219	protein phosphatase 2 (formerly 2A), regulatory subu
330	GTGACGCCTGT	1,00	0.61	Hs.279361	
	CCATAAGTCCT	1,00			seven in absentia (Drosophila) homolog
340	CCATAAGTCCT	1,00			1
341	AGCCCCGCGC	1,00	0,61	Hs.272320	Homo sapiens mRNA; cDNA DKFZp434L1226 (from clone DK
242	TATCATCATTC	1.00	0.64	Hs.270877	
	TTCCCTGAGCA	1,00			ESTs, Weakly similar to ALU1_HUMAN
343	TICCCIGAGCA	1,00	0,61	NS.20190	ALU SUBFAMILY J S
	TAGTCTAAGGC	1,00		Hs.261782	
	ATAGTAGTAAT	1,00		Hs.258863	
346	GATTTTAAATG	1,00			interleukin 1 receptor, type II
347	CCTGGCTAACA	1,00		Hs.252124	
348	CTGTACAGACC	1,00	0,61	Hs.251653	tubulin, beta, 2
349	GGCACCAGAGC	1,00		Hs.249614	
350	GCGAACTCCGT	1,00		Hs.248844	
351	CGCGTCCGTGT	1,00		Hs.243929	
352	GTTTTGGTTTA	1,00	0,61	Hs.241336	Homo sapiens mRNA; cDNA
					DKFZp564G0422 (from clone DK
	CTITTTGCCAC	1,00		Hs.240165	
354	GTGGGGGGCGC	1,00	0,61	Hs.240031	EST, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB S
355	TCTTTCCAACT	1,00	0.61	Hs.22394	hypothetical protein FLJ10893
	TAGTAGGGCTC	1,00		Hs.21914	
	AGGACATAACA	1,00		Hs.213793	
	TTGTTTTAAGA	1,00			hypothetical protein LOC56757
	GTGGCACATCT	1,00			ESTs, Weakly similar to alternatively
		.,	-,0 .		spliced produc
360	CACATTGAGGC	1,00	0.61	Hs.207122	
	AGGCTAGCACT	1,00			Homo sapiens mRNA for KIAA1190
					protein, partial cds
362	CCCCTGCCCTC	1,00	0,61	Hs.203317	EST

363 TTGTCTCTTGA 1,00 0,61 Hs.20104 [ESTs   364 TGGGTACACTG   1,00 0,61 Hs.200030 [ESTs, Weakly similar to ALUB_HUMAN   1918 GGCAGTGGTAA   1,00 0,61 Hs.197075 [ESTs   366 CATAAATGTTA   1,00 0,61 Hs.197075 [ESTs   367 AGATTACCCAC   1,00 0,61 Hs.192155 [ESTs   368 ACTGGCTCAGG   1,00 0,61 Hs.192155 [ESTs   369 CTTGTAGTCTC   1,00 0,61 Hs.190719 [ESTs   369 CTTGTAGTCTC   1,00 0,61 Hs.19073 [ESTs   370 AGCCCAGCTGG   1,00 0,61 Hs.18857 [ESTs   371 GAGGCCAACTG   1,00 0,61 Hs.18857 [ESTs   372 GAGCCAACTG   1,00 0,61 Hs.188675 [ESTs   373 GACCCAACCTGC   1,00 0,61 Hs.183601 [regulator of G-protein signalling 16   373 GCCCTACCTGC   1,00 0,61 Hs.182740 [ribosomal protein S11   374 GCCAACGGCGT   1,00 0,61 Hs.182740 [ribosomal protein S11   374 GCCAACGGCGT   1,00 0,61 Hs.182740 [ribosomal protein S11   377 GCGAACGAGAA   1,00 0,61 Hs.182740 [ribosomal protein S11   377 GCGAACGAGAA   1,00 0,61 Hs.175941 [B-cell receptor-associated protein BAP23   378 GCGAACAGGAA   1,00 0,61 Hs.175941 [B-cell receptor-associated protein BAP23   378 GGGGTAATTTT   1,00 0,61 Hs.167144 [ESTs, Highly similar to AF070470_1   379 CTGAAGTGCA   1,00 0,61 Hs.167144 [ESTs, Highly similar to AF070470_1   379 CTGAAGTGCAG   1,00 0,61 Hs.161554 [rypothetical protein FLJ20159   380 CCCATTCAGTC   1,00 0,61 Hs.161554 [rypothetical protein FLJ20159   381 TAATGTCTCA   1,00 0,61 Hs.161554 [rypothetical protein FLJ20159   383 GCAGATTCTCA   1,00 0,61 Hs.1516452 [ESTs   384 GGAATGAGGGG   1,00 0,61 Hs.152925 [KIAA1268 protein   1,00 0,61 Hs.153632 [ESTs   386 GAAGGAACTA   1,00 0,61 Hs.153632 [ESTs   388 GAGGAACTGAT   1,00 0,61 Hs.153632 [ESTs   388 GAGGAACTGA   1,00 0,61 Hs.153632 [ESTs   389 GAGGAACTA   1,00 0,61 Hs.153632 [ESTs   389 GAGGACCTGAA   1,00 0,61 Hs.13785 [ESTs   389 GAGGACCTGAA   1,00 0,61 Hs.13785 [ESTs   399 GAGGACCTGAA   1,00 0,61 Hs.138643 [ESTs   399 GAGGACCTGAA   1,00 0,61 Hs.138643 [ESTs   399 GAGGACCTGAA   1,00 0,61 Hs.						
IIII ALU CLASS B						
365   CATAAATGTTA	364	TGGGTACACTG	1,00			IIII ALU CLASS B
367   AGATTACCCAC	365	GGCAGTGGTAA	1,00	0,61	Hs.197075	ESTs
367   AGATTACCCAC	366	CATAAATGTTA	1,00	0,61	Hs.19479	ESTs
368   ACTGGCTCAGG	367	AGATTACCCAC	1,00	0,61	Hs.192155	ESTs
369   CTTGTAGTCTC						
370   AGCCCAGCTGG						
371   GAGGGCAATCT						
373   GCCCTACCTGC   1,00   0,61   Hs. 182740   ribosomal protein S11				0,61	Hs.186753	
373   GCCCTACCTGC   1,00   0,61   Hs. 182740   ribosomal protein S11	372	CTGAAACAGGA	1,00	0,61	Hs.183601	regulator of G-protein signalling 16
374   GCCAACGGCGT	373	GCCCTACCTGC				
376   GGGGTAATTTT   1,00   0,61   Hs.173497   Sec23 (S. cerevisiae)   homolog B	374	GCCAACGGCGT	1,00	0,61	Hs.181002	
377   AGGACAATGAA   1,00   0,61   Hs.173135   dual-specificity tyrosine-(Y)-phosphorylation regula   STR   TTTGCACTTTT   1,00   0,61   Hs.167114   ESTs, Highly similar to AF070470_1   SPARC-related pro   379   CTGAAGTGCAG   1,00   0,61   Hs.166609   ESTs   SPARC-related pro   381   TAATGTTCTCA   1,00   0,61   Hs.161554   hypothetical protein FLJ20159   381   TAATGTTCTCA   1,00   0,61   Hs.159642   glucosaminyl (N-acetyl) transferase 1, core 2 (beta-383   GCAGATTCTCA   1,00   0,61   Hs.157716   ESTs   384   GGAATGAGGGG   1,00   0,61   Hs.156452   ESTs   385   GAGAGGACTA   1,00   0,61   Hs.15323   ESTs   386   TTCGACAGCT   1,00   0,61   Hs.15323   ESTs   Weakly similar to ALUF_HUMAN   III   ALU CLASS F   389   GAGTATTATTT   1,00   0,61   Hs.144906   ESTs   STs   Weakly similar to EPS8_HUMAN   EPIDERMAL GROWTH   Sylvar	375	GTAAAGATGAA	1,00	0,61	Hs.175941	B-cell receptor-associated protein BAP29
phosphorylation regula   378 TTTGCACTTTT   1,00   0,61   Hs.167114   ESTs, Highly similar to AF070470_1   SPARC-related pro   379 CTGAAGTGCAG   1,00   0,61   Hs.166609   ESTs   380 CCCATTCAGTC   1,00   0,61   Hs.161554   hypothetical protein FLJ20159   381 TAATGTTCTCA   1,00   0,61   Hs.160271   G protein-coupled receptor 48   382 GTTTCTGCAGA   1,00   0,61   Hs.159642   glucosaminyl (N-acetyl) transferase 1, core 2 (beta-383 GCAGATTCTCA   1,00   0,61   Hs.157716   ESTs   384 GGAATGAGGGG   1,00   0,61   Hs.156452   ESTs   385 GAGAGGAACTA   1,00   0,61   Hs.15323   ESTs   386 TTCGACAGGCT   1,00   0,61   Hs.152925   KIAA1268 protein   387 CTCCAGCCTGG   1,00   0,61   Hs.144906   ESTs   389 GAGTATTATTT   1,00   0,61   Hs.144906   ESTs   Weakly similar to ALUF_HUMAN   IIII ALU CLASS F   Human   EPIDERMAL GROWTH   390 TACTGTACTCC   1,00   0,61   Hs.143198   hypothetical protein similar to tumor suppressor p33   AAAATAAAATG   1,00   0,61   Hs.142908   E2F-like protein   392 CCCAGGAGTTT   1,00   0,61   Hs.136433   ESTs   394 CATTTGGCCGG   1,00   0,61   Hs.136433   ESTs   395 CAGGACCTGAA   1,00   0,61   Hs.135260   ESTs   395 CAGGACCTGAA   1,00   0,61   Hs.135260   ESTs   397 CAGTAGGATAA   1,00   0,61   Hs.135251   ESTs   398 GCCTCACCTGG   1,00   0,61   Hs.135251   ESTs   399 TCTGTATCAAA   1,00   0,61   Hs.128814   ESTs   399 TCTGTACCAAA   1,00   0,61   Hs.128814   ESTs   399 TCTGTATCAAA   1,00   0,61   Hs.128814   ESTs   399 TCTGTATCAAA   1,00   0,61   Hs.128814   ESTs   399 TCTGTATCAAA   1,00   0,61   Hs.128820   ESTs   Moderately similar to   ALU1   HUMAN ALU SUBFAMILY   400 TGATTTGTGAA   1,00   0,61   Hs.12282   ESTs   STS	376	GGGGTAATTTT		0,61	Hs.173497	Sec23 (S. cerevisiae) homolog B
SPARC-related pro   379 CTGAAGTGCAG   1,00   0,61 Hs.166609 ESTs   380 CCCATTCAGTC   1,00   0,61 Hs.161554 hypothetical protein FLJ20159   381 TAATGTTCTCA   1,00   0,61 Hs.160271   G protein-coupled receptor 48   382 GTTTCTGCAGA   1,00   0,61 Hs.159642   glucosaminyl (N-acetyl) transferase 1, core 2 (beta-383 GCAGATTCTCA   1,00   0,61 Hs.157716 ESTs   384 GGAATGAGGGG   1,00   0,61 Hs.153623 ESTs   385 GAGAGGAACTA   1,00   0,61 Hs.153523 ESTs   386 TTCGACAGGCT   1,00   0,61 Hs.153523 ESTs   387 CTCCAGCCTGG   1,00   0,61 Hs.14331 ESTs, Weakly similar to ALUF_HUMAN	377	AGGACAATGAA	1,00			phosphorylation regula
380   CCCATTCAGTC   1,00   0,61   Hs.161554   hypothetical protein FLJ20159	378	TTTGCACTTTT	1,00	0,61	Hs.167114	
TAATGTTCTCA	379	CTGAAGTGCAG	1,00	0,61	Hs.166609	ESTs
382 GTTTCTGCAGA	380	CCCATTCAGTC	1,00	0,61	Hs.161554	hypothetical protein FLJ20159
Core 2 (beta-	381	TAATGTTCTCA	1,00	0,61	Hs.160271	G protein-coupled receptor 48
383   GCAGATTCTCA   1,00   0,61   Hs.157716   ESTs     384   GGAATGAGGGG   1,00   0,61   Hs.156452   ESTs     385   GAGAGGAACTA   1,00   0,61   Hs.153523   ESTs     386   TTCGACAGGCT   1,00   0,61   Hs.152925   KIAA1268 protein     387   CTCCAGCCTGG   1,00   0,61   Hs.145331   ESTs, Weakly similar to ALUF_HUMAN   IIII ALU CLASS F     389   GAGTATTATTT   1,00   0,61   Hs.144906   ESTs     389   GAGTATTATTT   1,00   0,61   Hs.143738   ESTs, Weakly similar to EPS8_HUMAN   EPIDERMAL GROWTH     390   TACTGTACTCC   1,00   0,61   Hs.143198   hypothetical protein similar to tumor   suppressor p33     391   AAAATAAAATG   1,00   0,61   Hs.142908   E2F-like protein     392   CCCAGGAGTTT   1,00   0,61   Hs.13785   ESTs     393   AAAATACAGTG   1,00   0,61   Hs.136433   ESTs     394   CATTTGGCCGG   1,00   0,61   Hs.136433   ESTs     395   CAGGACCTGAA   1,00   0,61   Hs.135260   ESTs     396   TCATTAACAAA   1,00   0,61   Hs.135260   ESTs     397   CAGTAGGATAA   1,00   0,61   Hs.134541   ESTs     398   GCCTCACCTGG   1,00   0,61   Hs.128514   ESTs     399   TCTGTATCAAA   1,00   0,61   Hs.128408   ESTs, Moderately similar to   ALU1_HUMAN ALU SUBFAMILY     400   TGATTTGTGAA   1,00   0,61   Hs.12282   ESTs	382	GTTTCTGCAGA	1,00	0,61	Hs.159642	
385   GAGAGGAACTA	383	GCAGATTCTCA	1,00	0,61	Hs.157716	ESTs
386         TTCGACAGGCT         1,00         0,61         Hs.152925         KIAA1268 protein           387         CTCCAGCCTGG         1,00         0,61         Hs.145331         ESTs, Weakly similar to ALUF_HUMAN IIII ALU CLASS F           388         AGTCTGTATTT         1,00         0,61         Hs.144906         ESTs           389         GAGTATTATTT         1,00         0,61         Hs.143738         ESTs, Weakly similar to EPS8_HUMAN EPIDERMAL GROWTH           390         TACTGTACTCC         1,00         0,61         Hs.143198         hypothetical protein similar to tumor suppressor p33           391         AAAATAAAATG         1,00         0,61         Hs.142908         E2F-like protein           392         CCCAGGAGTTT         1,00         0,61         Hs.13785         ESTs           393         AAAATACAGTG         1,00         0,61         Hs.136433         ESTs           394         CATTTGGCCGG         1,00         0,61         Hs.135971         ESTs           395         CAGGACCTGAA         1,00         0,61         Hs.135260         ESTs           397         CAGTAGGATAA         1,00         0,61         Hs.128514         ESTs           399         TCTGTATCAAA         1,00         0,61	384	GGAATGAGGGG	1,00	0,61	Hs.156452	ESTs
387         CTCCAGCCTGG         1,00         0,61         Hs.145331         ESTs, Weakly similar to ALUF_HUMAN IIII ALU CLASS F           388         AGTCTGTATTT         1,00         0,61         Hs.144906         ESTs           389         GAGTATTATTT         1,00         0,61         Hs.143738         ESTs, Weakly similar to EPS8_HUMAN EPIDERMAL GROWTH           390         TACTGTACTCC         1,00         0,61         Hs.143198         hypothetical protein similar to tumor suppressor p33           391         AAAATAAAATG         1,00         0,61         Hs.142908         E2F-like protein           392         CCCAGGAGTTT         1,00         0,61         Hs.13785         ESTs           393         AAAATACAGTG         1,00         0,61         Hs.136433         ESTs           394         CATTTGGCCGG         1,00         0,61         Hs.136031         ESTs           395         CAGGACCTGAA         1,00         0,61         Hs.135971         ESTs           396         TCATTAACAAA         1,00         0,61         Hs.134541         ESTs           398         GCCTCACCTGG         1,00         0,61         Hs.128514         ESTs           399         TCTGTATCAAA         1,00         0,61	385	GAGAGGAACTA	1,00	0,61	Hs.153523	ESTs
387         CTCCAGCCTGG         1,00         0,61         Hs.145331         ESTs, Weakly similar to ALUF_HUMAN IIII ALU CLASS F           388         AGTCTGTATTT         1,00         0,61         Hs.144906         ESTs           389         GAGTATTATTT         1,00         0,61         Hs.143738         ESTs, Weakly similar to EPS8_HUMAN EPIDERMAL GROWTH           390         TACTGTACTCC         1,00         0,61         Hs.143198         hypothetical protein similar to tumor suppressor p33           391         AAAATAAAATG         1,00         0,61         Hs.142908         E2F-like protein           392         CCCAGGAGTTT         1,00         0,61         Hs.13785         ESTs           393         AAAATACAGTG         1,00         0,61         Hs.136433         ESTs           394         CATTTGGCCGG         1,00         0,61         Hs.136031         ESTs           395         CAGGACCTGAA         1,00         0,61         Hs.135971         ESTs           396         TCATTAACAAA         1,00         0,61         Hs.134541         ESTs           398         GCCTCACCTGG         1,00         0,61         Hs.128514         ESTs           399         TCTGTATCAAA         1,00         0,61	386	TTCGACAGGCT	1,00	0,61	Hs.152925	KIAA1268 protein
389         GAGTATTATTT         1,00         0,61         Hs.143738         ESTs, Weakly similar to EPS8_HUMAN EPIDERMAL GROWTH           390         TACTGTACTCC         1,00         0,61         Hs.143198         hypothetical protein similar to tumor suppressor p33           391         AAAATAAAATG         1,00         0,61         Hs.142908         E2F-like protein           392         CCCAGGAGTTT         1,00         0,61         Hs.13785         ESTs           393         AAAATACAGTG         1,00         0,61         Hs.136433         ESTs           394         CATTTGGCCGG         1,00         0,61         Hs.136031         ESTs           395         CAGGACCTGAA         1,00         0,61         Hs.135971         ESTs           396         TCATTAACAAA         1,00         0,61         Hs.135260         ESTs           397         CAGTAGGATAA         1,00         0,61         Hs.128514         ESTs           399         TCTGTATCAAA         1,00         0,61         Hs.128408         ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           400         TGATTTGTGAA         1,00         0,61         Hs.12282         ESTs	387	CTCCAGCCTGG		0,61	Hs.145331	ESTs, Weakly similar to ALUF_HUMAN
EPIDERMAL GROWTH   390 TACTGTACTCC	388	AGTCTGTATTT	1,00	0,61	Hs.144906	ESTs
Suppressor p33	389	GAGTATTATTT	1,00	0,61	Hs.143738	
391 AAAATAAAATG         1,00         0,61 Hs.142908 E2F-like protein           392 CCCAGGAGTTT         1,00         0,61 Hs.13785 ESTs           393 AAAATACAGTG         1,00         0,61 Hs.136433 ESTs           394 CATTTGGCCGG         1,00         0,61 Hs.136031 ESTs           395 CAGGACCTGAA         1,00         0,61 Hs.135971 ESTs           396 TCATTAACAAA         1,00         0,61 Hs.135260 ESTs           397 CAGTAGGATAA         1,00         0,61 Hs.134541 ESTs           398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1 HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs	390	TACTGTACTCC	1,00	0,61		
393 AAAATACAGTG         1,00         0,61 Hs.136433 ESTs           394 CATTTGGCCGG         1,00         0,61 Hs.136031 ESTs           395 CAGGACCTGAA         1,00         0,61 Hs.135971 ESTs           396 TCATTAACAAA         1,00         0,61 Hs.135260 ESTs           397 CAGTAGGATAA         1,00         0,61 Hs.134541 ESTs           398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1 HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs	391	AAAATAAAATG	1,00	0,61		
394 CATTTGGCCGG         1,00         0,61 Hs.136031 ESTs           395 CAGGACCTGAA         1,00         0,61 Hs.135971 ESTs           396 TCATTAACAAA         1,00         0,61 Hs.135260 ESTs           397 CAGTAGGATAA         1,00         0,61 Hs.134541 ESTs           398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1 HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs	392	CCCAGGAGTTT	1,00	0,61	Hs.13785	ESTs
395 CAGGACCTGAA         1,00         0,61 Hs.135971 ESTs           396 TCATTAACAAA         1,00         0,61 Hs.135260 ESTs           397 CAGTAGGATAA         1,00         0,61 Hs.134541 ESTs           398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1 HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs			1,00			
396   TCATTAACAAA   1,00   0,61   Hs.135260   ESTs     397   CAGTAGGATAA   1,00   0,61   Hs.134541   ESTs     398   GCCTCACCTGG   1,00   0,61   Hs.128514   ESTs     399   TCTGTATCAAA   1,00   0,61   Hs.128408   ESTs, Moderately similar to   ALU1   HUMAN ALU SUBFAMILY   400   TGATTTGTGAA   1,00   0,61   Hs.12282   ESTs			1,00	0,61	Hs.136031	ESTs
397 CAGTAGGATAA         1,00         0,61 Hs.134541 ESTs           398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs	395	CAGGACCTGAA	1,00	0,61	Hs.135971	ESTs
398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs	396	TCATTAACAAA		0,61	Hs.135260	ESTs
398 GCCTCACCTGG         1,00         0,61 Hs.128514 ESTs           399 TCTGTATCAAA         1,00         0,61 Hs.128408 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           400 TGATTTGTGAA         1,00         0,61 Hs.12282 ESTs	397	CAGTAGGATAA				
399 TCTGTATCAAA 1,00 0,61 Hs.128408 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY 400 TGATTTGTGAA 1,00 0,61 Hs.12282 ESTs	398	GCCTCACCTGG				
400 TGATTTGTGAA 1,00 0,61 Hs.12282 ESTs	399	TCTGTATCAAA				ESTs, Moderately similar to
	400	TGATTTGTGAA	1,00	0,61	Hs.12282	
						<del></del>

400	070071010	1 4 00	0.04		1001.40
	GTGGTACACAG	1,00			CGI-43 protein
	TTGGCCAGATT	1,00			CGI-43 protein
	ATTCTTGTACA	1,00		Hs.117527	
	CCTTCTTGGGG	1,00		Hs.117474	
406	TACTGGAAGGC	1,00	0,61	Hs.116874	ESTs, Weakly similar to putative p150
					[H.sapiens]
	GTGGTGGTTGG	1,00			toll-like receptor 5
408	AGTAATGAAAA	1,00	0,61	Hs.11217	KIAA0877 protein
409	TTAGTTTTGCT	1,00	0,61	Hs.108885	collagen, type VI, alpha 1
410	GCTAATATATT	1,00	0,61	Hs.107883	ESTs
411	TCTTGACTCCC	1,00	0,61	Hs.107265	ESTs
	TGTACTTAATT	1,00			ESTs, Moderately similar to alternatively
		'	·		spliced pr
413	CTCAGCAGGAG	1,00	0.61	Hs.105489	ESTs, Weakly similar to AF109127_1
		'			stromal cell-deri
414	GGTACAATCCG	1,00	0.61	Hs.104557	hypothetical protein FLJ10697
	ACTCCAGACCT	1,00		Hs.104350	
-	GAGAAACTCCG	9,00			GTPase activating protein-like
	CCCAGAGACCC	19,00			calponin 1, basic, smooth muscle
	GCAAGAAAGTG	23,00	12.05	Hs 155376	hemoglobin, beta
-	AAACAATAAAA	8,00		Hs.229971	
	CGTGGGACACT	8,00			NICE-1 protein
	CTGTTCTCTTG	4,00		Hs.46824	
	TAGTTGGAAAA	33,00			nuclear receptor subfamily 4, group A,
722	17011007777	33,00	10,02	113.1119	member 1
423	GTGAAAGCCTG	6,00	3 35	Hs.258926	
	TTGGTTTGCTG	3,00			Human clone 23960 mRNA sequence
-	CCTGTAATTCA	3,00			EST, Weakly similar to ALU5_HUMAN
720	OCIGIAATIOA	3,00	1,70	115.277331	ALU SUBFAMILY SC S
126	AAACCCCGTCT	3,00	1 76	He 272464	ESTs, Weakly similar to ALU2_HUMAN
720	77700000101	3,00	1,70	115.213404	ALU SUBFAMILY SB
127	ATCGCACTACT	3,00	1 76	Hs.161721	
	TTGAGGGGGTG	17,00			(Manual assignment) MEMOREC
420	TIGAGGGGGTG	17,00	0,00	ns.70049	unassignable (probably r
120	TTTGGTTTTCC	115,00	56.20	Ho 170572	
$\overline{}$	ACAAAACCCCG	7,00		Hs.259505	collagen, type I, alpha 2
<del></del>		2,00			ESTs
	GTGCTCAATAG				
	CAACCAGTAAA	2,00		Hs.79914	
_	AGTTTATGCCC	2,00			KIAA0887 protein
	TTCACATTGTC	2,00		Hs.285804	
	CCTGGCCTAGA	2,00		Hs.285472	
	AAACTGGGAGG	2,00		Hs.231722	
	GCCGCCTTCT	2,00		Hs.201292	
	TGCAGGTTTGT	2,00			Ran GTPase activating protein 1
	CTCAACTTGTA	2,00			CGI-43 protein
	TTTGCTTTTGT	10,00			aquaporin 3
441	TAAATGAAAAA	3,00	1,69	Hs.82120	nuclear receptor subfamily 4, group A,
					member 2
442	GCCCCCTTCCT	3,00	1,69	Hs.212680	tumor necrosis factor receptor

	Τ	<del></del>	· <del>-</del> · · · · · · · · · · · · · · · · · · ·		superfamily, member 1
142	CCTCTAATTCC	200	4.00	115 404 404	
	CCTGTAATTGC	3,00		Hs.181464	
	TCACCCTCCAG	3,00			hypothetical protein
	AAGCTCTGTGT	5,00	2,67	Hs.19813	
	ATGGTGGGCGC	4,00		Hs.266417	
	CCTGTAGTTCT	4,00			ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC
448	CATCTGTAATC	4,00	2,19	Hs.153290	ESTs, Weakly similar to prostate-specific transgluta
449	TCTATAATCCC	6,00	3,14	Hs.96866	ESTs
450	ATGGCACGTGC	7,00			stromal cell protein
451	CTGGTGCACTG	2,00			ESTs, Weakly similar to ALU8_HUMAN ALU SUBFAMILY SX
452	ATAAAAAGAAA	2,00	1,11	Hs.83942	cathepsin K (pycnodysostosis)
453	CACTTGTAGTC	2,00		Hs.7845	Homo sapiens cDNA FLJ20820 fis, clone ADSE00490
	ATTTGGAGAGG	2,00			Fukuyama type congenital muscular dystrophy
455	CAGCAGCTTGT	2,00	1,11	Hs.55405	ESTs, Weakly similar to Rab7 [H.sapiens]
456	GTATTTTCATA	2,00	1,11	Hs.42140	hypothetical protein FLJ10103
457	CTTGTTGCAAT	2,00	1,11	Hs.29640	suppression of tumorigenicity 15 (reversion-inducing
458	CAGGGTGGGTG	2,00	1,11	Hs.278222	ESTs, Highly similar to endothelial nitric oxide syn
459	AGCCACTACGC	2,00	1.11	Hs.249956	
	TGGCATAATCA	2,00		Hs.237063	
	CCTTCCTCTCC	2,00		Hs.199752	
	CCCTGAATGAA	2,00			frizzled (Drosophila) homolog 4
	GATTAGCACCA	2,00			ribosomal protein L5
	GCCGTGAAAAA	2,00		Hs.164257	
	AGAAAGAAGGA	2,00		Hs.1501	syndecan 2 (heparan sulfate proteoglycan 1, cell sur
466	GCTTCCTCCTC	5,00	2.60		CD34 antigen
	ттстссст	5,00		Hs.283009	
-	TGCCTGTAGTC	111,00		Hs.285275	ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
469	CCCTCAATCCC	6,00	3,07	Hs.83077	interleukin 18 (interferon-gamma- inducing factor)
470	ACAACTTTTAT	6,00	3,07	Hs.283213	
471	GCAAACCTAAA	3,00	1,62	Hs.80686	prefoldin 5
	GAGTGCAACCC	3,00	1,62	Hs.54680	ESTs
473	TTGAATAGTGA	4,00	2,12	Hs.38516	ESTs
474	AGCCGGATGCT	4,00			KIAA0720 protein
475	GCAAAACACTG	4,00		_	Homo sapiens mRNA; cDNA DKFZp566B193 (from clone DKF
476	CATTTGGGAAG	4,00	2 12	Hs.111334	ferritin, light polypeptide
	CAAGAGATGCT	1,00			ESTs, Weakly similar to cell division
					control relate

478	AGTITATTTCA	1,00	0,54	Hs.99016	Human DNA sequence from clone 310J6 on chromosome 6q
479	GAGGATCTGCG	1,00	0.54	Hs.90998	KIAA0128 protein; septin 2
	ACTCTGGCTCA	1,00		Hs.88974	cytochrome b-245, beta polypeptide
.00	, 1010100010, 1	.,,,,	0,0 .		(chronic granulom
481	GGGGTTAGGGG	1,00	0.54	Hs.85050	phospholamban
	TAATATATCTG	1,00		Hs.8203	endomembrane protein emp70 precursor
				· · ·	isolog
483	TAAGTCTATAT	1,00	·	Hs.78864	Fc fragment of IgG, low affinity IIa, receptor for (
484	CACAAAAGGAT	1,00	0,54	Hs.77603	ESTs
485	GAGAAACCCTT	1,00	0,54	Hs.7739	ESTs
486	GAAAACAGTAA	1,00	0,54	Hs.76111	dystroglycan 1 (dystrophin-associated glycoprotein 1
487	GGCAATTTACT	1,00	0,54	Hs.75813	polycystic kidney disease 1 (autosomal dominant)
488	TTTTGTGCTAG	1,00	0,54	Hs.74649	cytochrome c oxidase subunit VIc
489	TTCCCGTGGCT	1,00		Hs.70983	PTPL1-associated RhoGAP 1
490	GGCCCAGGCCT	1,00		Hs.575	aldehyde dehydrogenase 3
	CGGCCACGTAT	1,00			ESTs
	TATAGCTGCAT	1,00		Hs.55964	ESTs, Weakly similar to C4HU complement C4A precurso
493	CTTGTGTTTAT	1,00	0.54	Hs.50748	chromosome 21 open reading frame 18
	CACAAAAATGC	1,00		Hs.49944	ESTs
	CTGTAATTTTA	1,00			ESTs
	CAATTCTTTCT	1,00			hypothetical protein FLJ10847
	ATTCTGCAGAG	1,00			ESTs
	TTTATATCATT	1,00			ESTs
	ACTCCATAAAA	1,00		Hs.4273	Human DNA sequence from clone RP1- 104A17 on chromoso
500	TTTTCATTATA	1,00	0,54	Hs.42656	ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
501	GACTCGACCAG	1,00	0,54	Hs.31922	ESTs
502	GGTGTGTTTTA	1,00	0,54	Hs.31566	ESTs
503	ATAAATTTATG	1,00	0,54	Hs.30715	ESTs
504	GGAGGCCGAGA	1,00	0,54	Hs.285565	ESTs
505	ATATTCAGCTG	1,00	0,54	Hs.285379	ESTs
506	GAACTTGTCTG	1,00	0,54	Hs.279934	Homo sapiens mRNA; cDNA DKFZp434I0835 (from clone DK
507	CCCGTATATGT	1,00	0.54	Hs.279844	hypothetical protein FLJ10033
	GTGGACCTGAG	1,00		Hs.279059	
	AGCCTGGAAGG	1,00		Hs.278549	
	TGGTTTTTGAG	1,00			ribosomal protein S18
				Hs.274969	
510		1.001	U. 341		
510 511	GCTTGTTCAAA	1,00			
510 511 512	GCTTGTTCAAA TCCACCAGCCA	1,00	0,54	Hs.27457	ESTs
510 511 512 513	GCTTGTTCAAA TCCACCAGCCA GCACTCCAACC	1,00 1,00	0,54 0,54	Hs.27457 Hs.273682	ESTs EST
510 511 512 513 514	GCTTGTTCAAA TCCACCAGCCA	1,00	0,54 0,54 0,54	Hs.27457 Hs.273682 Hs.270797	ESTs EST

517	TAAAGATGGCA	1,00	0,54	Hs.25357	Homo sapiens clone 24488 mRNA
					sequence
	GGCTATGCCCT	1,00		Hs.243855	
	AGACATTGACA	1,00			CGI-96 protein
	TATGCTTTAAA	1,00			ribosomal protein S5 pseudogene 1
521	TATTGCTAAAT	1,00	0,54	Hs.23590	solute carrier family 16 (monocarboxylic
					acid transp
	CCAAGGCACTG	1,00		Hs.234863	
	AACTTTCCAAA	1,00		Hs.23457	
	TCTCACAAGGG	1,00		Hs.233476	
525	CCACTATGCCT	1,00		Hs.231229	
526	GGGCGCCTGGC	1,00		Hs.224242	
527	GGGGGAAAAA	1,00			EST, Weakly similar to RL3_HUMAN 60S RIBOSOMAL PROTE
528	TTGTTTATGTA	1,00	0,54	Hs.21958	Homo sapiens cDNA FLJ10532 fis, clone NT2RP2001044
529	TTGAGATAAGA	1,00	0,54	Hs.21887	
530	GCTATATCCAA	1,00		Hs.218008	
531	TTTGGTTTTCT	1,00			suppressor of fused
	GATTGTCCTTG	1,00		Hs.211517	
	GGTTATCAAAG	1,00		Hs.208334	
	GTCTCGCTGAC	1,00		Hs.207911	
	AGCCATCGCGC	1,00		Hs.207749	
	ACCACCCGTGT	1,00		Hs.202033	
	ACACTCTTCCT	1,00		Hs.20103	
	GGAGATGTTTG	1,00		Hs.199545	
	CTCAACCTTAA	1,00			KIAA0231 protein
	GCAAGACTCCC	1,00		Hs.198011	
	CAATACTATTC	1,00			hypothetical protein FLJ10388
	GTTGTGCTCAG	1,00	0.54	Hs.191228	hypothetical protein FLJ20284
	GGGCTAGCACT	1,00	0.54	Hs.190722	FSTs
	ACATTCACGCC	1,00			glutaryl-Coenzyme A dehydrogenase
	TATATTTAGTT	1,00			protein kinase, cAMP-dependent,
0.,0		1,00	0,0-1	110.100001	regulatory, type I,
546	CCAGCTAGCGA	1,00	0.54	Hs.179756	
	TGAGGATACAG	1,00			Homo sapiens mRNA; cDNA
ا'``	TOROURINORO	1,00	0,54	113.177020	DKFZp564L102 (from clone DKF
548	TGCCTACAGTC	1,00	0.54	Hs.176207	
	CTCATATGCAA	1,00			ATPase, Class V, type 10D
	AAGGAGGTGGA	1,00		Hs.172730	
	CAAGCCCTGCC	1,00			hypothetical protein similar to mouse
		1,00	0,04	2000	HN1 (Hematolo
552	GCGCTGGGAGG	1,00	0.54	Hs 171763	CD22 antigen
	TCCTTGGCGTG	1,00			Ank, mouse, homolog of
	TTGCCATATGC	1,00			ESTs, Weakly similar to unnamed
554	ITTOOMINIO	1,00	0,54	110.104024	protein product [H.s
555	GCTCCCTTCAC	1,00	0.54	Hs.162222	
	CTGTGCAAGGA	1,00			hypothetical protein FLJ20159
22/	TGTGCTTGTGT	1,00	<u> </u>	MS. 161554	hypothetical protein FLJ20159

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558	СТТТТССССС	1,00	0.54	Hs.156007	Down syndrome critical region gene 1-
					like 1
	TCTTTTCTTT	1,00			paired mesoderm homeo box 1
	AAAAGCTGTTT	1,00		Hs.15550	
	GTCCCCCCCC	1,00		Hs.152454	
562	CCTGGCGGGAT	1,00	0,54	Hs.149347	ESTs, Highly similar to KIAA1043 protein [H.sapiens]
563	GATTGGTATGA	1,00	0,54	Hs.147049	cut (Drosophila)-like 1 (CCAAT
564	GTGAAGACTAC	1,00	0.54	Hs.14665	displacement protein) ESTs
	TAAATGGGTTG	1,00		Hs.1395	early growth response 2 (Krox-20
					(Drosophila) homolo
	TGTAAACTTTG	1,00		Hs.13849	
	GACACGTTGCC	1,00	0,54	Hs.136574	arachidonate 12-lipoxygenase, 12R type
	TGTGAATTTTA	1,00			hematopoietic protein 1
	TAAAACACTTG	1,00		Hs.130636	
	TGGCAATTTTC	1,00		Hs.129636	
571	TCTGTAGCACA	1,00			hypothetical protein FLJ10600
572	TCAGCAGTTAA	1,00	0,54	Hs.128571	ESTs
573	AATGAGGTGCT	1,00	0,54	Hs.128400	ESTs
574	TTCAGAATCTT	1,00	0,54	Hs.125914	ESTs
575	GAATAAGATAT	1,00	0,54	Hs.12479	associated molecule with the SH3 domain of STAM
576	TGTCCCAGCCA	1,00	0,54	Hs.1211	acid phosphatase 5, tartrate resistant
	CCCATCTCAGA	1,00		Hs.120769	Homo sapiens cDNA FLJ20463 fis, clone KAT06143
578	AGAAGCTCCAG	1,00	0,54	Hs.120021	DKFZP434I092 protein
579	AGTCTTCAAAA	1,00			CGI-43 protein
580	GAGTTTTCATT	1,00			CGI-43 protein
581	TTTTGCAATAA	1,00		Hs.11614	
-	AGCTCTATGAG	1,00		Hs.115831	
	AATAGGTCCCA	1,00			ribosomal protein S25
584	TAGTCTACTGT	1,00		Hs.112472	
	ATCCTTACATC	1,00		Hs.111720	
586	AACTTTCATAT	1,00			phosphatidylinositol glycan, class N
587	CCGTAAAAAAA	1,00			divalent cation tolerant protein CUTA
588	TGAAGATGTAA	1,00		Hs.106217	
$\overline{}$	GAACCATTTGC	1,00			KIAA0926 protein
	CTGTGGAGCTG	1,00		Hs.103379	
	AGAATTACAGA	1,00		Hs.101915	
	TGTTGCTCCCA	5,00			zinc finger protein 220
593	TAGTTTGAAGG	5,00			glutaminyl-peptide cyclotransferase (glutaminyl cycl
594	CCCAACGCGCT	47,00	20.67	Hs 272572	hemoglobin, alpha 2
	CCTATAATCTC	13,00			CGI-43 protein
	TCTCCTGGACT	4,00			kallikrein 5
	ATCTTGCCACT	4,00			glutamate receptor, metabotropic 1
	AGGATAAAAAA	3,00		Hs.79404	neuron-specific protein
	GACCACAAATA	3,00		Hs.76476	cathepsin H
	C. IOO/IOAATA	, 3,00	1,50	113.70470	oathepsiii i i

000	A COATATOTTO	0.00	4.50	11- 075005	leit and another 040
	AGCATATCTTC	3,00			ribosomal protein S18
601	AACACAGGAGG	3,00	1,56	HS.222874	ESTs, Moderately similar to zinc
000	00000100100		4.50	11 404040	transporter 4 [H.sa
602	CGGGGACGAGG	3,00	1,56	HS.124942	protein phosphatase 2A 48 kDa
-			- 1 1 1		regulatory subunit
603	GCGAAGCCCCG	7,00	3,35	Hs.103804	heterogeneous nuclear ribonucleoprotein
					U (scaffold
	GTGGCAGGTAC	9,00		Hs.190467	
	TCTGTTGTTCA	9,00			CGI-43 protein
	CCTGTATCCCA	4,00		Hs.270072	
607	ATGGATGCTTG	2,00	1,05	Hs.89404	msh (Drosophila) homeo box homolog 2
608	GACTTCTGTCC	2,00	1,05	Hs.87539	aldehyde dehydrogenase 8
609	AAAAAGAAACT	2,00	1,05	Hs.73287	KIAA1235 protein
610	TACTGAAAAAA	2,00	1,05	Hs.5111	hypothetical protein FLJ20729
611	CACCTGGAGGC	2,00	1,05		ESTs
612	GCAAGAGCCCA	2,00	1,05	Hs.26670	Human PAC clone RP3-515N1 from
1 1		٠ ]	·		22q11.2-q22
613	AACCCGGGGAG	2,00	1,05	Hs.228009	
	TTGCCCAGGGT	2,00			ESTs, Weakly similar to ALU5_HUMAN
1 1			·		ALU SUBFAMILY SC
615	AAGCAGTTACA	2,00	1.05	Hs.22116	CDC14 (cell division cycle 14, S.
			•		cerevisiae) homolo
616	GCAGTCATACA	2,00	1.05	Hs.182626	chromosome 22 open reading frame 5
$\overline{}$	GGGGCACACAC	2,00	1.05	Hs.181900	ESTs
$\rightarrow$	AGGGAAGGTGA	2,00		Hs.126927	
	AGCCGCTGTGC	2,00		Hs.106771	
	TCAAGCCATCA	35,00			early growth response 1
$\rightarrow$	GTGGCGGGCAT	5,00		Hs.230564	
	ACCTGGGTGCT	5,00			ESTs, Weakly similar to MLD [H.sapiens]
	TGTGGCGTATA	8,00			myosin, light polypeptide kinase
	CGGCACCTTC	6,00			gap junction protein, beta 5 (connexin
		-,	0,00		31.1)
625	CCCTTGAGGAG	6,00	8 99	Hs.1076	(Manual assignment) SPRR1B, cornifin
		3,55	0,00		B .
626	AATGTTTTTAA	3,00	1.50	Hs.75335	glycine amidinotransferase (L-
-		-,-5	.,		arginine:glycine amidi
627	CCTCTCCCATT	3,00	1.50	Hs.177533	Homo sapiens mRNA, chromosome 1
		-,	.,		specific transcript
628	TGTTCTGATTT	3,00	1.50		acyl-Coenzyme A oxidase 1, palmitoyl
	CCTGTAGTGCC	5,00			Homo sapiens cell-line E8CASS clone
		-,	_,,_,		E24L estradiol-i
630	CCACTGTATTC	6,00	2.77	Hs.235041	
	AGAAATGTATG	6,00			transcription factor 8 (represses
'''		5,55	~,' '		interleukin 2 expr
632	AGGTCAGAAGA	10,00	4 38	Hs.23437	Homo sapiens mRNA; cDNA
		.5,55	,,55	0.20701	DKFZp586G0623 (from clone DK
633	ATTAAGAAAAT	14,00	5.95	Hs.76549	AHNAK nucleoprotein (desmoyokin)
	AACCCGGGGGG	6,00		Hs.6214	KIAA0731 protein
	CACCACAACAA	6,00			chloride channel 3
	J. 100/10/VI	0,00	۲,12	1.3.11-113	omonde charmer 3

620	TTCCCCACCCT	2.00	0.00	U. 50545	Line finesy protoin 45
	TTCCCCAGGGT	2,00			ring finger protein 15
637	CACACTATAGG	2,00	0,99	Hs.58924	ESTs, Weakly similar to JC5594 jerky
600	00000000	2 00	0.00	11- 25000	gene protein ho
$\overline{}$	GGGAAAGAGGG	2,00			KIAA1538 protein
	GCTGGGCGCGG	2,00		Hs.278070	
$\overline{}$	GGGGCAACAGC	2,00			CDW52 antigen (CAMPATH-1 antigen)
	GTGGCAGGCCC	2,00		Hs.266105	
	CATATCCCCTC	2,00		Hs.250746	
	CTTAGGAGTCA	2,00		Hs.23853	
	CAGCACAGTGG	2,00			ras GTPase activating protein-like
645	ATGCTCAAAGG	2,00			Homo sapiens mRNA full length insert cDNA clone EURO
646	TTCTGTGCATA	2,00			hypothetical protein FLJ10231
647	GCATAATGTTT	2,00		Hs.11050	F-box only protein 9
648	TGGCCAGCTCC	19,00	7,83		protein tyrosine phosphatase, receptor type, C
649	TCTGGCCCAGC	3,00	1,45	Hs.183	Duffy blood group
650	ATCCTGAGTTA	12,00			major histocompatibility complex, class II, DQ beta
651	ACAAATGAAAA	1,00	0,49	Hs.96657	hyothetical protein
	TACATTTCAAG	1,00		Hs.94376	proprotein convertase subtilisin/kexin type 5
653	GCAAAATGCTG	1,00	0.49		hypothetical protein FLJ20163
	TTTAAGAAATG	1,00			ESTs
	ACCCACCTGTG	1,00			Homo sapiens mRNA; cDNA DKFZp564H203 (from clone DKF
656	GCAGCAGTGTC	1,00	0.49	Hs.86538	ESTs
	CCATTAAAAAA	1,00			ESTs
~	AAAGAGGGACG	1,00		Hs.84229	splicing factor, arginine/serine-rich 8 (suppressor-
659	ACACTTAAAAA	1,00	0.49	Hs.83381	guanine nucleotide binding protein 11
	TCACAAAAAA	1,00		Hs.7976	KIAA0332 protein
	ATTTAATATAT	1,00		Hs.7972	KIAA0871 protein
	AGTTCAAGGCT	1,00		Hs.79058	suppressor of Ty (S.cerevisiae) 4 homolog 1
663	GAGTTGCTATT	1,00	0,49	Hs.78575	prosaposin (variant Gaucher disease and variant meta
664	GACTGTCAAAA	1,00	0,49	Hs.69330	ESTs
	GCGCAACAACT	1,00		Hs.61950	DKFZp434A0131 protein
	TAAATCAGAGA	1,00		Hs.61929	Homo sapiens cDNA FLJ11010 fis, clone PLACE1003145
667	TAAATGAATGA	1,00	0,49	Hs.57967	ESTs
	AATAGGAAGAT	1,00			ESTs
669	CAATTAAAAAT	1,00	0,49	Hs.50123	zinc finger protein 189
	GTGTATGTGGT	1,00			fatty acid binding protein 3, muscle and
			,		heart (mamm
671	ACAAACCCACA	1,00	0.49	Hs.49282	hypothetical protein FLJ11088
	CAAGTGAAAGG	1,00		Hs.47822	KIAA0380 gene product; RhoA-specific
		.,	-, · •		guanine nucleot

070	04404044000	4 661	0.40	11- 47500	FOT-
	GAACAGAACGC	1,00		Hs.47566	
	CTAATGGCCCT	1,00		Hs.3416	adipose differentiation-related protein
	TAATGTTTTT	1,00			ESTs
676	ATTTGGCCTGT	1,00	0,49	Hs.285519	Homo sapiens OVN6-2 mRNA, partial
					cds
	TCTGGGAGGGG	1,00	0,49	Hs.285313	core promoter element binding protein
	GACTAAATTGT	1,00		Hs.283643	
	TATTTTCACAA	1,00			hypothetical protein FLJ20546
680	GAGGTTTTCTG	1,00	0,49	Hs.279639	Homo sapiens mRNA; cDNA
					DKFZp586M2022 (from clone DK
681	CCTAGAATCCC	1,00			PRO1779 protein
682	CAGCCCCTGTC	1,00	0,49	Hs.278234	Homo sapiens mRNA; cDNA
					DKFZp434H1323 (from clone DK
683	GTGTCGGGCTC	1,00	0,49	Hs.278010	EST
684	CACCTGTAAAC	1,00	0,49	Hs.277311	EST
685	GCGGCGACTGC	1,00	0,49	Hs.271980	mitogen-activated protein kinase 6
686	GGAATGAATGA	1,00		Hs.270824	
	TGGGATATAGT	1,00	0,49	Hs.269888	ESTs
	GTGGTTGATGC	1,00			ESTs, Weakly similar to ALU7_HUMAN
		.,	-,		ALU SUBFAMILY SQ
689	GTGGTAGGTAC	1,00	0.49	Hs.264844	EST, Moderately similar to
		.,	٠, ٠٠		ALU1_HUMAN ALU SUBFAMILY
690	GCCTGCCTTTA	1,00	0.49	Hs.25371	ESTs, Weakly similar to A37232 mucin,
		.,	-,		tracheal [H.sa
691	GATCTCGCTTT	1,00	0.49	Hs.250773	signal sequence receptor, alpha
•••		.,00	0, .0	. 10.200. 10	(translocon-associat
692	CTCCTATTTTT	1,00	0.49	Hs 25010	hypothetical protein P15-2
	GTGCGCACCTG	1,00			ESTs, Weakly similar to ALU1_HUMAN
333		.,00	٥, .٠		ALU SUBFAMILY J S
694	ATAAAATGTCT	1,00	0.49	Hs.24181	ESTs
	TTTCACCCCGT	1,00			ESTs
	TATGTTTAAAA	1,00			platelet-activating factor acetylhydrolase
555		1,00	0,40	110.204002	2 (40kD)
697	GATCACTTGAG	1,00	0.49	Hs.231798	
	TTGAGAAAAA	1,00		Hs.22971	
-	ACCCTGGGAGG	1,00		Hs.228529	
	TGTGGTGGTGC	1,00		Hs.223618	
	GATTGGCCAGA	1,00			
'''	GATTGGCCAGA	1,00	0,48	115.21/39	Homo sapiens mRNA; cDNA DKFZp586I1518 (from clone DK
702	GCTGAGTTATT	1,00	0.40	Hs.216363	
	AGTATTCCTAA	1,00			hypothetical protein DKFZp762O076
-	TGTCTGCCATT			Hs.21580	
	CCCCTGTACTC	1,00			
(05	COCCIGIACIO	1,00	0,49	⊓S.∠13016	ESTs, Highly similar to cytokine receptor
700	CTITITAAAAA	4.00	0.40	11- 040700	related pr
	CTTTTTAAAGA	1,00		Hs.212788	
'''	TACCCGAAAAC	1,00	0,49	ms.210858	ESTs, Weakly similar to ALUD_HUMAN
700	TOOTOTAGE	4.56	0.46	LI- 000440	IIII ALU CLASS D
108	TGCCTCTAGTC	1,00	0,49	ms.209413	ESTs, Weakly similar to ALU1_HUMAN
					ALU SUBFAMILY J S

700	000000000	4 00	0.40	11- 004040	Irox.
	CCGCCGCACTC	1,00		Hs.204049	
	CTAAGAGGGGT	1,00		Hs.196102	
	CTGCTGTAGTC	1,00		Hs.194319	
	GGACCCTCATT	1,00		Hs.191063	
	AACTAATTCTC	1,00	0,49	Hs.189513	hypothetical protein FLJ10213
714	GCAGTGCCAAG	1,00	0,49	Hs.183528	ESTs, Weakly similar to Bem46-like
					protein [D.melano
	ATTGGAGCGCA	1,00			rTS beta protein
	TGAAAACTCCC	1,00	0,49	Hs.180248	zinc finger protein 124 (HZF-16)
717	CTCCCAGCCAC	1,00	0,49	Hs.177582	surfactant, pulmonary-associated protein
					A1
	GTTTAAAAAA	1,00	0,49	Hs.174756	EST
	GAATCAAGCTG	1,00		Hs.17296	
720	CTGGGCATTTT	1,00	0,49	Hs.172207	non-POU-domain-containing, octamer-
					binding
	TTGTTTGTGTA	1,00			KIAA1357 protein
	AGGACTGGCAT	1,00			KIAA0061 protein
	TCGCTTTTAAG	1,00			Ank, mouse, homolog of
	AAGTGAAAAAA	1,00	0,49	Hs.168159	apoptosis regulator
725	CAAATTAGAAT	1,00	0,49		ESTs, Moderately similar to
					CO3_HUMAN COMPLEMENT C3
726	ATACCACTAAG	1,00	0,49	Hs.153792	5-methyltetrahydrofolate-homocysteine
	·			<u> </u>	methyltransfer
727	GGTGATGGAGG	1,00	0,49	Hs.149692	ESTs, Highly similar to G43284 zinc
					finger protein Z
	TTGGGTTTCTG	1,00		Hs.147975	
729	AATAAATGCCC	1,00		Hs.145522	
730	CATACACACAT	1,00			protein kinase C, nu
	GTTATACAACA	1,00		Hs.141183	
	CTCACAGGCAC	1,00		Hs.139784	
733	TACCCATTACC	· 1,00		Hs.136981	
734	TTTCTTTCCCT	1,00		Hs.135055	
735	CATCTGTACTG	1,00	0,49	Hs.132892	Homo sapiens protocadherin 10
					(PCDH10) mRNA, partial
-	GAGCTGTTTTG	1,00		Hs.127476	
737	ACACGTACTAT	1,00	0,49	Hs.120828	Human DNA sequence from clone RP5-
	•		•		876B10 on chromoso
738	TTTATATTTCA	1,00	0,49	Hs.11958	oxidative 3 alpha hydroxysteroid
					dehydrogenase; reti
	CACAGTTTTAA	1,00		Hs.118918	
740	TGGTTTTACCA	1,00			CGI-43 protein
	GTGCCTGGTAT	1,00			hypothetical protein PRO2198
742	TACCTTAGAAC	1,00	0,49	Hs.114963	Homo Sapiens (clone B3B3E13)
					chromosome 4p16.3 DNA f
743	GTGTAGGAGGT	1,00	0,49	Hs.113029	ribosomal protein S25
	CCCTGGAGACA	1,00	0,49	Hs.111334	ferritin, light polypeptide
745	CCTGTAGCCCA	1,00	0,49	Hs.109370	ESTs
746	TATCCATATTA	1,00	0,49	Hs.107942	DKFZP564M112 protein
747	CTACTAATTGC	1,00	0,49	Hs.101916	Homo sapiens mRNA; cDNA

					DKFZp564K133 (from clone DKF
	GGCCTCCAAGA	5,00		Hs.286220	
749	GTGGCTGACAC	4,00			ESTs, Weakly similar to ALU4_HUMAN ALU SUBFAMILY SB2
750	CTGTACTTGTG	8,00	3,37	Hs.75678	FBJ murine osteosarcoma viral oncogene homolog B
751	ACAAAACCCCA	8,00	3,37	Hs.140208	
	AGCCACCACCC	3,00			ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
753	TGCCTGTAGTT	7,00	2,97	Hs.246646	EST
754	GCTAACCCCTG	15,00	5,92	Hs.279772	brain specific protein
755	GCAAAACCCTG	86,00			CGI-43 protein
756	AAACATTAAAA	21,00			actin, gamma 2, smooth muscle, enteric
757	GTGGCACTTGC	4,00	1,78	Hs.272322	Homo sapiens mRNA; cDNA DKFZp434L092 (from clone DKF
758	GAAGCTACACC	4,00		Hs.107253	
759	TAAGGTAGAGG	2,00	0,94	Hs.99908	nuclear receptor coactivator 4
760	TGGGTCATTTG	2,00	0,94	Hs.98073	ESTs
761	GCTCACTGAAG	2,00	0,94	Hs.9568	zinc finger protein 261
762	CACCTATCAAT	2,00	0,94	Hs.58617	Rho-associated, coiled-coil containing protein kinas
763	CTTCAATCTTA	2,00	0.94	Hs.58419	DKFZP586L2024 protein
	CAGTCCTCTTG	2,00			tousled-like kinase 2
	TTACCAAAGCA	2,00		Hs.30246	solute carrier family 19 (thiamine transporter), mem
766	ATGAAACCCTA	2,00	0.94	Hs.282671	
	TCACTCCAGCC	2,00		Hs.270497	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
768	AAAGGCATCAG	2,00	0.94		integrin, alpha 11
	GACATCTGTCC	2,00		Hs.25566	
	TGTCTTTTCTG	2,00			bradykinin receptor B2
	TGACTGTATTA	2,00			amine oxidase, copper containing 3 (vascular adhesio
772	GCAAGACCTCA	2,00	0,94	Hs.181592	ESTs
773	AGGAAGGAAAA	2,00		Hs.180532	heat shock 90kD protein 1, alpha
774	CTGCCGGAGCA	2,00	0,94	Hs.164779	ESTs
775	TATTCCAGAAC	2,00			hypothetical protein FLJ20159
776	AGCGAAACTCC	2,00		Hs.106597	
777	CTGGGGGTCAG	2,00			vesicle-associated membrane protein 4
778	GCTGTAATCCC	15,00	5,82	Hs.184019	Homo sapiens clone 23551 mRNA sequence
779	CACCTGTGGTC	16,00	6,11	Hs.209585	EST
780	CTCTAGAGAAA	3,00			hypothetical protein
781	GCAAATCCTGT	3,00		Hs.79059	transforming growth factor, beta receptor III (betag
782	TATATGCTGGG	3,00	1,35	Hs.2969	v-ski avian sarcoma viral oncogene homolog
783	GGGAGACCCCA	3,00	1,35	Hs.24884	ESTs, Moderately similar to RNA polymerase I associa

784	TGGAAATGAAA	3,00	1 35	He 172028	collagen, type I, alpha 1
	CCTGTAATCTG	3,00		Hs.159975	
	TGTATTGTACA	3,00			Link guanine nucleotide exchange factor
					11
787	ACATAGACCGA	7,00			pigment epithelium-derived factor
788	GTTCCACAGAA	12,00	4,66		collagen, type I, alpha 2
	GTGGCAGAGAC	4,00			polycystic kidney disease 1 (autosomal dominant)
	AGCCGAGATCG	4,00		Hs.278053	
	CCTGTAGTTCC	24,00	8,84	Hs.189242	Homo sapiens mRNA; cDNA DKFZp434A202 (from clone DKF
792	GTAAAACCCTG	29,00	10,50	Hs.281680	peroxisomal trans 2-enoyl CoA reductase; putative sh
793	AGTCTGCTGGG	5,00	7,44	Hs.259508	
	CTGGGCAGAGA	5,00		Hs.200735	ESTs, Moderately similar to AF105377_1 heparan sulfa
795	TCACACAAAGG	5,00	2,11	Hs.46783	<del></del>
	CGCCTGTAATT	5,00		Hs.14333	
797	ATTGCTCTCTG	8,00		Hs.245188	tissue inhibitor of metalloproteinase 3 (Sorsby fund
798	CCTGTGGTTCC	10,00	3.88	Hs.286061	
	TGTCCACACAT	4,00			Homo sapiens mRNA; cDNA DKFZp586P1622 (from clone DK
800	GACGGAGCCTT	4,00	1,69	Hs.37482	COPZ2 for nonclathrin coat protein zeta- COP
801	ACTGGGCAGTG	4,00	1,69	Hs.241257	latent transforming growth factor beta binding prote
802	CGGCACCTTAA	4,00	1,69	Hs.209100	DKFZP434C171 protein
803	CTGAAACAGCT	4,00			suppressor of var1 (S.cerevisiae) 3-like 1
804	TGTTACCTGGT	2,00		Hs.79227	myomesin (M-protein) 2 (165kD)
805	TCTTTGCTCTT	2,00	0,90	Hs.44077	hypothetical protein FLJ10793
806	AAATCACCAAT	2,00		Hs.4082	lectin, galactoside-binding, soluble, 8 (galectin 8)
807	TGTATTGACTG	2,00	0,90	Hs.29280	ESTs
	TTCACTTCAAC	2,00	0,90		Homo sapiens clone 23967 unknown mRNA, partial cds
	CGTGGGGCTGC	2,00	0,90	Hs.221986	aquaporin 5
	TCACAGAGTCT	2,00	0,90	Hs.194149	Homo sapiens mRNA; cDNA DKFZp434M011 (from clone DKF
811	TGCCACTGTGC	2,00	0,90	Hs.183165	ESTs, Highly similar to protein kinase [H.sapiens]
812	CTATAAAAGTG	2,00	0,90	Hs.154145	guanine nucleotide binding protein (G protein), alph
813	TTGGCTAGGCT	13,00	4.86	Hs.252259	ribosomal protein S3
	AGGGAGCAGAG	11,00			microfibrillar-associated protein 4
	TTGGTGTGCTG	7,00		Hs.240399	
	TGGAAAGTGAA	50,00			v-fos FBJ murine osteosarcoma viral oncogene homolog
817	TTGATTGAGTG	3,00	1 21	Hs.9879	ESTs
	· · · · · · · · · · · · · · · · · · ·	3,00	1,01	10.3013	LU 13

818	CAGGGATCTGC	3,00	1,31	Hs.7634	ESTs, Moderately similar to semaphorin B [M.musculus
819	ATTGCACCACC	3,00	1,31	Hs.44259	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
820	CTGGTGGCCAC	3,00	1,31	Hs.286028	Human alpha-2 collagen type VI mRNA, 3' end
821	AATCATTGAGG	3,00			Homo sapiens mRNA; cDNA DKFZp564A2164 (from clone DK
822	ACCCCCTTCCT	3,00			major histocompatibility complex, class I, E
823	CCTGGCCAAAA	3,00		Hs.126824	
824	TTAACCCCTCC	30,00			ribonuclease, RNase A family, 1 (pancreatic)
825	ACAGGCTACGG	118,00	39,00	Hs.75777	transgelin
826	AGATGAGATGA	17,00	6,08	Hs.285313	core promoter element binding protein
827	AAAAAAAAGGC	1,00	0,44	Hs.90077	TG-interacting factor (TALE family homeobox)
828	TACGCTAAAAC	1,00	0,44	Hs.87354	ESTs
829	GTAGAAGTGTA	1,00	0,44	Hs.8705	ESTs
830	TGCCCAGCAAT	1,00	0,44	Hs.76297	G protein-coupled receptor kinase 6
831	AGCAGGTTTGC	1,00	0,44	Hs.7434	ESTs
	AGAGAGAGCCC	1,00			thyroid hormone receptor, alpha (avian erythroblasti
833	GCACTGATTAA	1,00	0,44		ESTs, Highly similar to I38945 melanoma ubiquitous m
834	CAGAGACAAGC	1,00	0,44	Hs.71721	ESTs
835	CAGTTGTCTAG	1,00	0,44	Hs.58882	Microfibril-associated glycoprotein-2
836	TGTGTGTGCTA	1,00	0,44	Hs.55533	ESTs
837	GCCTGGACCAG	1,00	0,44	Hs.55407	Homo sapiens mRNA; cDNA DKFZp434K0621 (from clone DK
838	TGTTTAATAAA	1,00	0,44	Hs.50841	ESTs
839	CATTITCTAAT	1,00	0,44	Hs.48376	Homo sapiens clone HB-2 mRNA sequence
840	AGCTTTCCCAA	1,00	0,44	Hs.45109	ESTs
841	ACTCAGTAGCC	1,00	0,44	Hs.44197	hypothetical protein DKFZp564D0462
	TCAAATTGAAA	1,00		Hs.44038	pellino (Drosophila) homolog 2
843	GGCCAGCCCTG	1,00	0,44	Hs.4243	ESTs
844	ACATTTTGTTC	1,00			ESTs
	GGCAAGCAGGC	1,00	0,44	Hs.41688	dual specificity phosphatase 8
846	TGCTTGGTACA	1,00	0,44	Hs.393	prostaglandin I2 (prostacyclin) receptor (IP)
847	TAAGTCTAATT	1,00		Hs.35804	hect domain and RLD 3
	TATTTTACTTG	1,00			hypothetical protein KIAA1165
849	CTCTCTCCCAG	1,00	0,44	Hs.30172	ESTs
850	AGCAAATTTTC	1,00	0,44	Hs.29423	ESTs, Weakly similar to macrophage lectin 2 [H.sapie
851	CCAAGACCTCT	1,00	0,44	Hs.283619	zinc finger protein 236
852	GACAGGTTCTG	1,00			transforming, acidic coiled-coil containing protein

	TTCTGGAGAA	1,00			hypothetical protein
854 T	GGGTTTTGTT	1,00	0,44	Hs.26243	Homo sapiens cDNA FLJ11177 fis, clone PLACE1007402
855 C	CACCGCTGCAG	1,00	0,44	Hs.261373	adenosine A2b receptor pseudogene
<del></del>	BAAACAGGAAA	1,00		Hs.257387	
	CTTCTTGCTA	1,00			hypothetical protein FLJ20171
	TAATCTGAAG	1,00			KIAA0022 gene product
	TAAATATAGG	1,00			leptin receptor gene-related protein
	ATGCTATGGT	1,00			ESTs
	STGAATGAAAC	1,00		Hs.223437	EST
862 T	CCACAAAAA	1,00			KIAA1130 protein
	GAATTGCTTA	1,00		Hs.203188	
864 C	CTCTAATTCC	1,00			ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
865 T	TGGACAAGAA	1,00		Hs.189902	<del></del>
	GCTGGGCGCG	1,00		Hs.188339	
	CAACACAAAGC	1,00			ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
868 G	GCCCGGCCTC	1,00	0,44	Hs.183994	protein phosphatase 1, catalytic subunit, alpha isof
	CTAAAAAATT	1,00			glucose regulated protein, 58kD
	TTTCCTTGTT	1,00	0,44	Hs.182937	peptidylprolyl isomerase A (cyclophilin A)
871 G	CCGCCTCTGT	1,00		Hs.180799	ESTs, Moderately similar to ALU5_HUMAN ALU SUBFAMILY
	GCTGGTTCCA	1,00	0,44		ribosomal protein L11
873 A	AGGAAATGAA	1,00			ribosomal protein L6
874 A	ATAGATGATA	1,00	0,44	Hs.174104	ESTs
875 T	ATAAGTGGAC	1,00	0,44	Hs.17301	ESTs
	BAAAACATAAT	1,00			DKFZP564A043 protein
	TTTGTAAAAA	1,00		Hs.16578	
	GCTGTTTAAA	1,00		Hs.164480	
	GGGACAGAGC	1,00	0,44	Hs.161554	hypothetical protein FLJ20159
	TAAAGCCAAA	1,00			ZAP3 protein
	CCCTGCATTC	1,00			chromosome 1 open reading frame 1
882 G	CTGAGAAGCA	1,00	0,44		protein tyrosine phosphatase, receptor type, C-assoc
883 G	BAATACGTTGG	1,00	0,44		BCL2/adenovirus E1B 19kD-interacting protein 2
884 A	TTTCTAACAA	1,00	0,44	Hs.155049	hypothetical protein FLJ11282
	GATCTTCTTG	1,00		Hs.14894	trans-Golgi network protein (46, 48, 51kD isoforms)
886 A	AAAACTCTAC	1,00	0,44	Hs.146226	
887 A	GGATAAACTC	1,00		Hs.14427	
888 C	TTGTGAGGCC	1,00			KIAA1161 protein
889 G	TAACAGTAAT	1,00		Hs.137396	
	AAACGAAAAT	1,00	0,44	Hs.135465	ESTs
891 A	GGGGAATGGG	1,00	0,44	Hs.134933	ESTs
	GCCGTGCTGC	1,00	0,44	Hs.131034	ESTs
893 G	GAACCTATCC	1,00	0,44	Hs.128807	ESTs

B95  AGCCAAGAGCC		0040700070	4 001	0.44	11. 400000	FOX
B95   AGCCAAGAGCC			1,00			
B97  GTAGACTGAAA						
898   TGGTATGCACC	-					
B89   TCTGTTTTGTG						
900   CCAATCAATGG						
OO1   TTCCCCAGGCT						
902   CAACAGCCCA						
903   CTCCTGGAATA   1,00	-					
905   TGGTGAAGAAC						
905   CACACCGCCCG	_					
			_			
906   CCCCCCCCA	905	CACACCGCCCG	1,00	0,44		
DKFZp434A139 (from clone DKF						[H.sapiens
907 TTTTGTTGCTC         1,00         0,44 Hs.109641 ESTs           908 GTAAAACCCCG         26,00         9,00 Hs.258881 EST, Weakly similar to ALU8_HUMAN ALU SUBFAMILY SX S           909 TAATAAAGAAT         4,00         1,65 Hs.80342 keratin 15           910 AAAATAAACCT         4,00         1,65 Hs.74304 periplakin           911 GTGGTAGGTGC         4,00         1,65 Hs.254237 EST           912 GTGAAATTCCA         4,00         1,65 Hs.258168 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           913 CCTCTAGTCCC         4,00         1,65 Hs.145501 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           914 TGGAAATGACC         106,00         34,28 Hs.172928 collagen, type I, alpha 1           915 AACCCTGGAGG         7,00         2,69 Hs.287355 ESTs           917 CCACTGCATTG         3,00         1,26 Hs.278551 ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918 AACTGGCTGCT         3,00         1,26 Hs.173381 dihydropyrimidinase-like 2           919 CCTGTAACCCC         28,00         9,54 Hs.176541 ESTs           920 GCACTCCAGCC         32,00         10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK St.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK St.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK St.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK St.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK St.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK St.	906	CCCCCCCCCA	1,00	0,44	Hs.110953	Homo sapiens mRNA; cDNA
908   GTAAAACCCCG   26,00   9,00   Hs.258881   EST, Weakly similar to ALU8_HUMAN ALU SUBFAMILY SX S     909   TAATAAAGAAT   4,00   1,65   Hs.80342   keratin 15     910   AAAATAAACCT   4,00   1,65   Hs.74304   periplakin     911   GTGGTAGGTGC   4,00   1,65   Hs.254237   EST     912   GTGAAATTCCA   4,00   1,65   Hs.228168   ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S     913   CCTCTAGTCCC   4,00   1,65   Hs.145501   ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S     914   TGGAAATGACC   106,00   34,28   Hs.172928   collagen, type I, alpha 1     915   AACCCTGGAGG   7,00   2,69   Hs.283927   Homo sapiens cytomegalovirus partial fusion receptor     916   AGGATCACTTG   3,00   1,26   Hs.278551   ESTs   Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB     918   AACTGGCTGCT   3,00   1,26   Hs.173381   dihydropyrimidinase-like 2     919   CCTGTAACCCC   28,00   9,54   Hs.176541   ESTs     920   GCACTCCAGCC   32,00   1,97   Hs.27311   single-minded (Drosophila) homolog 2     921   GTGGTGAGCAC   5,00   1,97   Hs.27311   single-minded (Drosophila) homolog 2     922   ATGGCAGGTGC   16,00   5,61   Hs.236479   EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S     923   CCATTACACTC   8,00   3,00   Hs.8594   KIAA1191 protein     924   GGTGACAGAGT   12,00   4,30   Hs.278314   hypothetical protein     925   GTGGTTTGCTG   9,00   3,31   Hs.157773   ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV     926   TCTGCCTATGC   4,00   1,60   Hs.90291   laminin, beta 2 (laminin S)     927   GCGAAAACCCC   4,00   1,60   Hs.95834   ESTs     928   TACCCCTTGAA   2,00   0,86   Hs.60548   hypothetical protein     929   TAAATGGAAGT   2,00   0,86   Hs.60548   hypothetical protein     920   TGTGCTCTGGGA   2,00   0,86   Hs.60548   hypothetical protein						
ALU SUBFAMILY SX S						
909   TAATAAAGAAT	908	GTAAAACCCCG	26,00	9,00		
910   AAAATAAACCT   4,00   1,65   Hs.74304   periplakin     911   GTGGTAGGTGC   4,00   1,65   Hs.254237   EST     912   GTGAAATTCCA   4,00   1,65   Hs.228168   ESTs, Weakly similar to ALU1_HUMAN     ALU SUBFAMILY J S     913   CCTCTAGTCCC   4,00   1,65   Hs.145501   ESTs, Weakly similar to ALU1_HUMAN     ALU SUBFAMILY J S     914   TGGAAATGACC   106,00   34,28   Hs.172928   collagen, type I, alpha 1     915   AACCCTGGAGG   7,00   2,69   Hs.283927   Homo sapiens cytomegalovirus partial   fusion receptor     916   AGGATCACTTG   3,00   1,26   Hs.287355   ESTs     917   CCACTGCATTG   3,00   1,26   Hs.278551   ESTs, Weakly similar to ALU2_HUMAN     ALU SUBFAMILY SB     918   AACTGGCTGCT   3,00   1,26   Hs.173381   dihydropyrimidinase-like 2     919   CCTGTAACCCC   28,00   9,54   Hs.176541   ESTs     920   GCACTCCAGCC   32,00   10,69   Hs.274580   Homo sapiens mRNA; cDNA     DKFZP434P1019 (from clone DK     921   GTGGTGAGCAC   5,00   1,97   Hs.27311   single-minded (Drosophila) homolog 2     922   ATGGCAGGTGC   16,00   5,61   Hs.236479   EST, Weakly similar to ALU7_HUMAN     923   CCATTACACTC   8,00   3,00   Hs.8594   KIAA1191 protein     924   GGTGACAGAGT   12,00   4,30   Hs.278314   hypothetical protein     925   GTGGTTTGCTG   9,00   3,31   Hs.157773   ESTs, Moderately similar to     926   TCTGCCTATGC   4,00   1,60   Hs.90291   alminin, beta 2 (laminin S)     927   GCGAAAACCCC   4,00   1,60   Hs.90291   laminin, beta 2 (laminin S)     928   TACCCCTTGAA   2,00   0,86   Hs.95834   ESTs     929   TAAATGGAAGT   2,00   0,86   Hs.950548   hypothetical protein     920   TGTGCTGGGA   2,00   0,86   Hs.60548   hypothetical protein     920   TGTGCTGGGA   2,00   0,86   Hs.60548   hypothetical protein						
911 GTGGTAGGTGC         4,00         1,65 Hs.254237 EST           912 GTGAAATTCCA         4,00         1,65 Hs.228168 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           913 CCTCTAGTCCC         4,00         1,65 Hs.145501 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           914 TGGAAATGACC         106,00         34,28 Hs.172928 collagen, type I, alpha 1           915 AACCCTGGAGG         7,00         2,69 Hs.283927 Homo sapiens cytomegalovirus partial fusion receptor           916 AGGATCACTTG         3,00         1,26 Hs.287355 ESTs           917 CCACTGCATTG         3,00         1,26 Hs.278551 ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918 AACTGGCTGCT         3,00         1,26 Hs.173381 dihydropyrimidinase-like 2           919 CCTGTAACCCC         28,00         9,54 Hs.176541 ESTs           920 GCACTCCAGCC         32,00         10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZP434P1019 (from clone DK SINGLE ST)           921 GTGGTGAGCAC         5,00         1,97 Hs.27311 single-minded (Drosophila) homolog 2           922 ATGGCAGGTGC         16,00         5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594 KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.1						
912         GTGAAATTCCA         4,00         1,65         Hs.228168         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           913         CCTCTAGTCCC         4,00         1,65         Hs.145501         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           914         TGGAAATGACC         106,00         34,28         Hs.172928         collagen, type I, alpha 1           915         AACCCTGGAGG         7,00         2,69         Hs.283927         Homo sapiens cytomegalovirus partial fusion receptor           916         AGGATCACTTG         3,00         1,26         Hs.287355         ESTs           917         CCACTGCATTG         3,00         1,26         Hs.278551         ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918         AACTGGCTGCT         3,00         1,26         Hs.173381         dihydropyrimidinase-like 2           919         CCTGTAACCCC         28,00         9,54         Hs.176541         ESTs           920         GCACTCCAGCC         32,00         10,69         Hs.274580         Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921         GTGGTGAGCAC         5,00         1,97         Hs.27341         single-minded (Drosophila) homolog 2           922         ATGGCAGGTGC         16,00         5,61	<del></del>					
Substitution   Subs						
ALU SUBFAMILY J S	912	GTGAAATTCCA	4,00	1,65	Hs.228168	
914         TGGAAATGACC         106,00         34,28         Hs.172928         collagen, type I, alpha 1           915         AACCCTGGAGG         7,00         2,69         Hs.283927         Homo sapiens cytomegalovirus partial fusion receptor           916         AGGATCACTTG         3,00         1,26         Hs.287355         ESTs           917         CCACTGCATTG         3,00         1,26         Hs.278551         ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918         AACTGGCTGCT         3,00         1,26         Hs.173381         dihydropyrimidinase-like 2           919         CCTGTAACCCC         28,00         9,54         Hs.176541         ESTs           920         GCACTCCAGCC         32,00         10,69         Hs.274580         Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921         GTGGTGAGCAC         5,00         1,97         Hs.27311         single-minded (Drosophila) homolog 2           922         ATGGCAGGTGC         16,00         5,61         Hs.236479         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923         CCATTACACTC         8,00         3,00         Hs.8594         KIAA1191 protein           924         GGTGACAGAGT         12,00         4,30         Hs.278314 hypothetical pro	913	CCTCTAGTCCC	4,00	1,65	Hs.145501	
915         AACCCTGGAGG         7,00         2,69         Hs.283927         Homo sapiens cytomegalovirus partial fusion receptor           916         AGGATCACTTG         3,00         1,26         Hs.287355         ESTs           917         CCACTGCATTG         3,00         1,26         Hs.278551         ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918         AACTGGCTGCT         3,00         1,26         Hs.173381         dihydropyrimidinase-like 2           919         CCTGTAACCCC         28,00         9,54         Hs.176541         ESTs           920         GCACTCCAGCC         32,00         10,69         Hs.274580         Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921         GTGGTGAGCAC         5,00         1,97         Hs.27311         single-minded (Drosophila) homolog 2           922         ATGGCAGGTGC         16,00         5,61         Hs.236479         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923         CCATTACACTC         8,00         3,00         Hs.8594         KIAA1191 protein           924         GGTGACAGAGT         12,00         4,30         Hs.157773         ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926         TCTGCCTATGC         4,00         1,60         Hs	914	TGGAAATGACC	106.00	34.28	Hs.172928	
Susion receptor   916 AGGATCACTTG   3,00   1,26 Hs.287355 ESTs   917 CCACTGCATTG   3,00   1,26 Hs.278551 ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB   918 AACTGGCTGCT   3,00   1,26 Hs.173381 dihydropyrimidinase-like 2   919 CCTGTAACCCC   28,00   9,54 Hs.176541 ESTs   920 GCACTCCAGCC   32,00   10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZP434P1019 (from clone DK   921 GTGGTGAGCAC   5,00   1,97 Hs.27311   single-minded (Drosophila) homolog 2   922 ATGGCAGGTGC   16,00   5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S   923 CCATTACACTC   8,00   3,00 Hs.8594   KIAA1191 protein   924 GGTGACAGAGT   12,00   4,30 Hs.278314   hypothetical protein   925 GTGGTTTGCTG   9,00   3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV   926 TCTGCCTATGC   4,00   1,60 Hs.90291   laminin, beta 2 (laminin S)   927 GCGAAAACCCC   4,00   1,60 Hs.272837 ESTs   928 TACCCCTTGAA   2,00   0,86 Hs.95834   ESTs   929 TAAATGGAAGT   2,00   0,86 Hs.6942   ESTs   930 TGTGTCTGGGA   2,00   0,86 Hs.60548   hypothetical protein PRO1635						
916 AGGATCACTTG         3,00         1,26 Hs.287355 ESTs           917 CCACTGCATTG         3,00         1,26 Hs.278551 ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918 AACTGGCTGCT         3,00         1,26 Hs.173381 dihydropyrimidinase-like 2           919 CCTGTAACCCC         28,00         9,54 Hs.176541 ESTs           920 GCACTCCAGCC         32,00         10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921 GTGGTGAGCAC         5,00         1,97 Hs.27311 single-minded (Drosophila) homolog 2           922 ATGGCAGGTGC         16,00         5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594 KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.6942 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.60548 hypothetical protein PRO1635	• • •	, , , , , , , , , , , , , , , , , , , ,	',""	-,		
917         CCACTGCATTG         3,00         1,26         Hs.278551         ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB           918         AACTGGCTGCT         3,00         1,26         Hs.173381         dihydropyrimidinase-like 2           919         CCTGTAACCCC         28,00         9,54         Hs.176541         ESTs           920         GCACTCCAGCC         32,00         10,69         Hs.274580         Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921         GTGGTGAGCAC         5,00         1,97         Hs.27311         single-minded (Drosophila) homolog 2           922         ATGGCAGGTGC         16,00         5,61         Hs.236479         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923         CCATTACACTC         8,00         3,00         Hs.8594         KIAA1191 protein           924         GGTGACAGAGT         12,00         4,30         Hs.278314         hypothetical protein           925         GTGGTTTGCTG         9,00         3,31         Hs.157773         ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926         TCTGCCTATGC         4,00         1,60         Hs.272837         ESTs           928         TACCCCTTGAA         2,00         0,86         Hs.6942         ESTs	916	AGGATCACTTG	3.00	1,26		
918 AACTGGCTGCT         3,00         1,26 Hs.173381 dihydropyrimidinase-like 2           919 CCTGTAACCCC         28,00         9,54 Hs.176541 ESTs           920 GCACTCCAGCC         32,00         10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921 GTGGTGAGCAC         5,00         1,97 Hs.27311 single-minded (Drosophila) homolog 2           922 ATGGCAGGTGC         16,00         5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594 KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635						ESTs, Weakly similar to ALU2_HUMAN
919 CCTGTAACCCC         28,00         9,54 Hs.176541 ESTs           920 GCACTCCAGCC         32,00         10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921 GTGGTGAGCAC         5,00         1,97 Hs.27311 single-minded (Drosophila) homolog 2           922 ATGGCAGGTGC         16,00         5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594 KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635	918	AACTGGCTGCT	3.00	1.26	Hs.173381	
920 GCACTCCAGCC         32,00         10,69 Hs.274580 Homo sapiens mRNA; cDNA DKFZp434P1019 (from clone DK           921 GTGGTGAGCAC         5,00         1,97 Hs.27311 single-minded (Drosophila) homolog 2           922 ATGGCAGGTGC         16,00         5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594 KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635						
DKFZp434P1019 (from clone DK						
921 GTGGTGAGCAC         5,00         1,97 Hs.27311         single-minded (Drosophila) homolog 2           922 ATGGCAGGTGC         16,00         5,61 Hs.236479         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594         KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635	5-5		32,33	.0,00	1 10121	
922 ATGGCAGGTGC         16,00         5,61 Hs.236479 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           923 CCATTACACTC         8,00         3,00 Hs.8594 KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635	921	GTGGTGAGCAC	5.00	1.97	Hs.27311	
ALU SUBFAMILY SQ S						
923 CCATTACACTC         8,00         3,00 Hs.8594         KIAA1191 protein           924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635				.,		
924 GGTGACAGAGT         12,00         4,30 Hs.278314 hypothetical protein           925 GTGGTTTGCTG         9,00         3,31 Hs.157773 ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926 TCTGCCTATGC         4,00         1,60 Hs.90291 laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635	923	CCATTACACTC	8.00	3.00	Hs.8594	
925         GTGGTTTGCTG         9,00         3,31         Hs.157773         ESTs, Moderately similar to OPSB_HUMAN BLUE-SENSITIV           926         TCTGCCTATGC         4,00         1,60         Hs.90291         laminin, beta 2 (laminin S)           927         GCGAAAACCCC         4,00         1,60         Hs.272837         ESTs           928         TACCCCTTGAA         2,00         0,86         Hs.95834         ESTs           929         TAAATGGAAGT         2,00         0,86         Hs.6942         ESTs           930         TGTGTCTGGGA         2,00         0,86         Hs.60548         hypothetical protein PRO1635						
OPSB_HUMAN BLUE-SENSITIV						
926 TCTGCCTATGC         4,00         1,60 Hs.90291   laminin, beta 2 (laminin S)           927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548   hypothetical protein PRO1635			'	-,		
927 GCGAAAACCCC         4,00         1,60 Hs.272837 ESTs           928 TACCCCTTGAA         2,00         0,86 Hs.95834 ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942 ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548 hypothetical protein PRO1635	926	TCTGCCTATGC	4,00	1,60	Hs.90291	
928 TACCCCTTGAA         2,00         0,86 Hs.95834         ESTs           929 TAAATGGAAGT         2,00         0,86 Hs.6942         ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548         hypothetical protein PRO1635	$\overline{}$					
929 TAAATGGAAGT         2,00         0,86 Hs.6942         ESTs           930 TGTGTCTGGGA         2,00         0,86 Hs.60548         hypothetical protein PRO1635						
930 TGTGTCTGGGA 2,00 0,86 Hs.60548 hypothetical protein PRO1635						<del></del>
						<del></del>
, 90 1100 1000 1 2,001 0.00103.04020 1E313		GCTTCAGTGGG	2,00		Hs.54828	ESTs
932 CTTTACTGTGT 2,00 0,86 Hs.279853 HSPC018 protein						

933   CACTTTGTGTG   2,00   0,86   Hs.24752   spectrin SH3 domain binding protein 1
935   ACTAAGATTGA   2,00   0,86   Hs.227716   KIAA0934 protein   936   CCAATGTTGTT   2,00   0,86   Hs.22209   ESTs   STs   0,86   Hs.194637   Homo sapiens mRNA; cDNA DKFZp564D113 (from clone DKF   938   TGAAGAATGTG   2,00   0,86   Hs.112557   ESTs, Moderately similar to ZN85   HUMAN ZINC FINGER P   939   GCCACTGCCTG   2,00   0,86   Hs.106061   RD RNA-binding protein   940   CCACTGCTCTC   12,00   4,26   Hs.23510   Kruppel-like factor 12   941   CTATTGCACTC   6,00   2,29   Hs.160483   erythrocyte membrane protein band 7.2 (stomatin)   epithelial membrane protein band 7.2 (stomatin)   942   ATGTACCTGAT   7,00   2,61   Hs.29191   epithelial membrane protein 2   943   GTGGGGGGGAG   7,00   2,61   Hs.29191   epithelial membrane protein 2   945   TCTGCACACAC   3,00   1,22   Hs.78518   natriuretic peptide receptor B/guanylate cyclase B (
936   CCAATGTTGTT   2,00   0,86   Hs.22209   ESTs     937   GTGGTGGAGC   2,00   0,86   Hs.194637   Homo sapiens mRNA; cDNA     938   TGAAGAATGTG   2,00   0,86   Hs.112557   ESTs, Moderately similar to     939   GCCAGTGCCTG   2,00   0,86   Hs.106061   RD RNA-binding protein     940   CCACTGCTCTC   12,00   4,26   Hs.23510   Kruppel-like factor 12     941   CTATTGCACTC   6,00   2,29   Hs.160483   erythrocyte membrane protein band 7.2 (stomatin)     942   ATGTACCTGAT   7,00   2,61   Hs.10700   hypothetical protein     943   GTGGGGGGGAG   7,00   2,61   Hs.10700   hypothetical protein     944   CACCTGTAGTC   37,00   11,92   Hs.267812   sorting nexin 4     945   TCTGCACACAC   3,00   1,22   Hs.78518   natriuretic peptide receptor B/guanylate cyclase B (     946   GACAATTCTGT   3,00   1,22   Hs.185571   hypothetical protein FLJ10700     947   AGGGGAAGGTG   8,00   2,92   Hs.112540   EST     948   CAAGACGGGGG   19,00   6,29   Hs.106185   ral guanine nucleotide dissociation stimulator     949   TAATTTGCGTT   4,00   1,56   Hs.79368   epithelial membrane protein 1     950   AGAATCGTTTG   4,00   1,56   Hs.268728   ESTs     951   AACAGTCAAAA   17,00   5,66   Hs.268728   ESTs     952   GCAAAACTCTG   10,00   3,49   Hs.278746   ESTs   Moderately similar to     ALUB   HUMAN ALU SUBFAMILY   S     953   GCCAGCCAGTG   10,00   3,49   Hs.149098   smoothelin     954   GCGAAATCCCA   6,00   2,21   Hs.268728   ESTs     955   TCTGTGGTCC   6,00   2,21   Hs.227894   ESTs     956   CCTATAAATTC   6,00   2,21   Hs.227894   ESTs     957   GCAATAAATAA   1,00   0,40   Hs.99651   ESTs     958   TGGAATCCAGG   1,00   0,40   Hs.99658   Homo sapiens clone 25023 mRNA     960   TTATTGTATTG   1,00   0,40   Hs.89474   ADP-ribosylation factor 6     961   ATGGAGGTATG   1,00   0,40   Hs.89444   ADP-ribosylation factor 6     961   ATGGAGGTATG   1,00   0,40   Hs.89444   ADP-ribosylation factor 6     961   ATGGAGGTATG   1,00   0,40   Hs.89444   ADP-ribosylation factor 6     962   ATGGAGGTATG   1,00   0,40   Hs.89444   ADP-ribosylation factor 6     963   ATGGAGG
937 GTGGTGTGAGC         2,00         0,86 Hs.194637 DKFZp564D113 (from clone DKF DKFZp564D113 (from clone DKF DKFZp564D113 (from clone DKF DKFZp564D113 (from clone DKF STS, Moderately similar to ZN85_HUMAN ZINC FINGER P           939 GCCAGTGCCTG         2,00         0,86 Hs.112557 ESTs, Moderately similar to ZN85_HUMAN ZINC FINGER P           940 CCACTGCTCTC         12,00         4,26 Hs.23510 Kruppel-like factor 12           941 CTATTGCACTC         6,00         2,29 Hs.160483 erythrocyte membrane protein band 7.2 (stomatin)           942 ATGTACCTGAT         7,00         2,61 Hs.29191 epithelial membrane protein 2 hypothetical protein           943 GTGGGGGGAG         7,00         2,61 Hs.10700 hypothetical protein           944 CACCTGTAGTC         37,00         1,92 Hs.267812 sorting nexin 4           945 TCTGCACACAC         3,00         1,22 Hs.186571 hypothetical protein FLJ10700           947 AGGGGAAGGTG         8,00         2,92 Hs.106185 ral guanine nucleotide dissociation stimulator           948 CAAGACGGGGG         19,00         6,29 Hs.106185 ral guanine nucleotide dissociation stimulator           949 TAATTTGCGTT         4,00         1,56 Hs.26557 plakophilin 3           951 AACAGTCAAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALUB_HUMAN ALU SUBFAMILY           953 GCAGCAGCG         6,00         2,21 Hs
DKFZp564D113 (from clone DKF
938   TGAAGAATGTG   2,00   0,86   Hs.112557   ESTs, Moderately similar to ZN85   HUMAN ZINC FINGER P     939   GCCAGTGCTCTC   12,00   4,26   Hs.23510   Kruppel-like factor 12     941   CTATTGCACTC   6,00   2,29   Hs.160483   erythrocyte membrane protein band 7.2 (stomatin)     942   ATGTACCTGAT   7,00   2,61   Hs.29191   epithelial membrane protein 2     943   GTGGGGGGGAG   7,00   2,61   Hs.10700   hypothetical protein     944   CACCTGTAGTC   37,00   11,92   Hs.267812   sorting nexin 4     945   TCTGCACACAC   3,00   1,22   Hs.186571   hypothetical protein FLJ10700     946   GACAATTCTGT   3,00   1,22   Hs.186571   hypothetical protein FLJ10700     947   AGGGGAAGGTG   8,00   2,92   Hs.112540   EST     948   CAAGACGGGGG   19,00   6,29   Hs.106185   ral guanine nucleotide dissociation   stimulator     949   TAATTTGCGTT   4,00   1,56   Hs.79368   epithelial membrane protein 1     950   AGAATCGTTTG   4,00   1,56   Hs.136299   ESTs     951   AACAGTCAAAA   17,00   5,66   Hs.26557   plakophilin 3     952   GCAAAACTCTG   10,00   3,49   Hs.278746   ESTs, Moderately similar to   ALUB   HUMAN ALU SUBFAMILY     953   GCCAGCCAGTG   10,00   3,49   Hs.149098   smoothelin     954   GCGAAATCCCA   6,00   2,21   Hs.268728   ESTs     955   TCTGTGGTCCC   6,00   2,21   Hs.27894   ESTs, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J S     956   CCTATAAATTCC   6,00   2,21   Hs.135491   ESTs     957   GCAATAAAATA   1,00   0,40   Hs.99621   ESTs     958   TGGGATCCAGG   1,00   0,40   Hs.99858   Homo sapiens clone 25023 mRNA   sequence     960   TTATTGTATTG   1,00   0,40   Hs.89474   ADP-ribosylation factor 6     961   ATGGAGGTATG   1,00   0,40   Hs.89474   ADP-ribosylation factor 6
Substitution
939   GCCAGTGCCTG   2,00   0,86   Hs. 106061   RD RNA-binding protein
940   CCACTGCTCTC   12,00   4,26   Hs.23510   Kruppel-like factor 12   941   CTATTGCACTC   6,00   2,29   Hs.160483   erythrocyte membrane protein band 7.2 (stomatin)   942   ATGTACCTGAT   7,00   2,61   Hs.29191   epithelial membrane protein 2   943   GTGGGGGGGAG   7,00   2,61   Hs.10700   hypothetical protein   944   CACCTGTAGTC   37,00   11,92   Hs.267812   sorting nexin 4   945   TCTGCACACAC   3,00   1,22   Hs.186571   hypothetical protein FLJ10700   947   AGGGGAAGGTG   8,00   2,92   Hs.112540   EST   948   CAAGACGGGG   19,00   6,29   Hs.106185   ral guanine nucleotide dissociation   949   TAATTTGCGTT   4,00   1,56   Hs.79368   epithelial membrane protein 1   950   AGAATCGTTTG   4,00   1,56   Hs.136299   ESTs   951   AACAGTCAAAA   17,00   5,66   Hs.26557   plakophilin 3   952   GCAAAACTCTG   10,00   3,49   Hs.278746   ESTs, Moderately similar to   ALUB   HUMAN   ALU   SUBFAMILY   953   GCCAGCCAGTG   10,00   3,49   Hs.149098   smoothelin   954   GCGAAATCCCA   6,00   2,21   Hs.268728   ESTs   955   TCTGTGGTCC   6,00   2,21   Hs.268728   ESTs   955   TCTGTGGTCCC   6,00   2,21   Hs.268728   ESTs   956   CCTATAATTCC   6,00   2,21   Hs.268728   ESTs   957   GCAATAAAATA   1,00   0,40   Hs.99621   ESTs   958   TGGAATCCAGG   1,00   0,40   Hs.99621   ESTs   959   CTTTTGTCAGC   1,00   0,40   Hs.99858   Homo sapiens clone 25023 mRNA   960   TTATTGTATTG   1,00   0,40   Hs.89474   ADP-ribosylation factor 6   961   ATGGAGGTATG   1,00   0,40   Hs.89474   ADP-ribosylation factor 6   961   ATGGAGGTATG   1,00   0,40   Hs.89474   ADP-ribosylation factor 6
941         CTATTGCACTC         6,00         2,29         Hs.160483         erythrocyte membrane protein band 7.2 (stomatin)           942         ATGTACCTGAT         7,00         2,61         Hs.29191         epithelial membrane protein 2           943         GTGGGGGGAG         7,00         2,61         Hs.10700         hypothetical protein           944         CACCTGTAGTC         37,00         11,92         Hs.78518         natriuretic peptide receptor B/guanylate cyclase B (           945         TCTGCACACAC         3,00         1,22         Hs.186571         hypothetical protein FLJ10700           947         AGGGGAAGGTG         8,00         2,92         Hs.112540         EST           948         CAAGACGGGGG         19,00         6,29         Hs.106185         ral guanine nucleotide dissociation stimulator           949         TAATTTGCGTT         4,00         1,56         Hs.79368         epithelial membrane protein 1           950         AGAATCGTTTG         4,00         1,56         Hs.136299         ESTs           951         AACAGTCAAAA         17,00         5,66         Hs.26572         plakophilin 3           952         GCAAGACCGGG         10,00         3,49         Hs.278746         ESTs, Moderately similar to ALU8 HUMAN ALU SUBFAMI
(stomatin)   (stomatin)   (pithelial membrane protein 2   (pithelial membrane protein 3   (pithelial membrane protein 4   (pithelial membrane protein 3   (pithelial membrane protein 3   (pithelial membrane protein 3   (pithelial membrane protein 5   (pithelial membrane protein 5   (pithelial membrane protein 5   (pithelial membrane protein 6   (pithelial membrane protein 1   (p
943         GTGGGGGGAG         7,00         2,61 Hs.10700         hypothetical protein           944         CACCTGTAGTC         37,00         11,92 Hs.267812         sorting nexin 4           945         TCTGCACACAC         3,00         1,22 Hs.78518         natriuretic peptide receptor B/guanylate cyclase B (           946         GACAATTCTGT         3,00         1,22 Hs.186571         hypothetical protein FLJ10700           947         AGGGGAAGGTG         8,00         2,92 Hs.112540         EST           948         CAAGACGGGGG         19,00         6,29 Hs.106185         ral guanine nucleotide dissociation stimulator           949         TAATTTGCGTT         4,00         1,56 Hs.79368         epithelial membrane protein 1           950         AGAATCGTTTG         4,00         1,56 Hs.136299         ESTs           951         JACAGTCAAAA         17,00         5,66 Hs.26557         plakophilin 3           952         GCAAAACTCTG         10,00         3,49 Hs.149098         smoothelin           953         GCCAGCCAGTG         10,00         3,49 Hs.278746         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY           955         TCTGTGGTCCC         6,00         2,21 Hs.268728         ESTs           955         TCTGTGGTCCC         6,00
944 CACCTGTAGTC         37,00         11,92 Hs.267812 sorting nexin 4           945 TCTGCACACAC         3,00         1,22 Hs.78518 natriuretic peptide receptor B/guanylate cyclase B (           946 GACAATTCTGT         3,00         1,22 Hs.186571 hypothetical protein FLJ10700           947 AGGGGAAGGTG         8,00         2,92 Hs.112540 EST           948 CAAGACGGGGG         19,00         6,29 Hs.106185 ral guanine nucleotide dissociation stimulator           949 TAATTTGCGTT         4,00         1,56 Hs.79368 epithelial membrane protein 1           950 AGAATCGTTTG         4,00         1,56 Hs.136299 ESTs           951 AACAGTCAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.99621 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.998135 ESTs
945         TCTGCACACAC         3,00         1,22         Hs.78518         natriuretic peptide receptor B/guanylate cyclase B (           946         GACAATTCTGT         3,00         1,22         Hs.186571         hypothetical protein FLJ10700           947         AGGGGAAGGTG         8,00         2,92         Hs.112540         EST           948         CAAGACGGGGG         19,00         6,29         Hs.106185         ral guanine nucleotide dissociation stimulator           949         TAATTTGCGTT         4,00         1,56         Hs.79368         epithelial membrane protein 1           950         AGAATCGTTTG         4,00         1,56         Hs.136299         ESTs           951         AACAGTCAAAA         17,00         5,66         Hs.26577         plakophilin 3           952         GCAAAACTCTG         10,00         3,49         Hs.278746         ESTs, Moderately similar to ALUS BFAMILY           953         GCCAGCCAGTG         10,00         3,49         Hs.149098         smoothelin           954         GCGAAATCCCA         6,00         2,21         Hs.268728         ESTs           955         TCTGTGGTCCC         6,00         2,21         Hs.135491         ESTs           957         GCAATAAAATA
Cyclase B (   946 GACAATTCTGT   3,00   1,22 Hs.186571 hypothetical protein FLJ10700   947 AGGGGAAGGTG   8,00   2,92 Hs.112540 EST   948 CAAGACGGGGG   19,00   6,29 Hs.106185 ral guanine nucleotide dissociation stimulator   949 TAATTTGCGTT   4,00   1,56 Hs.79368   epithelial membrane protein 1   950 AGAATCGTTTG   4,00   1,56 Hs.136299 ESTs   951 AACAGTCAAAA   17,00   5,66 Hs.26557   plakophilin 3   952 GCAAAACTCTG   10,00   3,49 Hs.278746   ESTs, Moderately similar to ALUB_HUMAN ALU SUBFAMILY   953 GCCAGCCAGTG   10,00   3,49 Hs.149098   smoothelin   954 GCGAAATCCCA   6,00   2,21 Hs.268728   ESTs   955 TCTGTGGTCCC   6,00   2,21 Hs.227894   ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S   956 CCTATAATTCC   6,00   2,21 Hs.135491   ESTs   957 GCAATAAAATA   1,00   0,40 Hs.99621   ESTs   958 TGGAATCCAGG   1,00   0,40 Hs.99135   ESTs   959 CTTTTGTCAGC   1,00   0,40 Hs.99135   ESTs   959 CTTTTGTCAGC   1,00   0,40 Hs.9944   ADP-ribosylation factor 6   961 ATGGAGGTATG   1,00   0,40 Hs.89474   ADP-ribosylation factor 6   961 ATGGAGGTATG   1,00   0,40 Hs.89444   procollagen C-endopeptidase enhancer   2
946 GACAATTCTGT         3,00         1,22 Hs.186571 hypothetical protein FLJ10700           947 AGGGGAAGGTG         8,00         2,92 Hs.112540 EST           948 CAAGACGGGG         19,00         6,29 Hs.106185 ral guanine nucleotide dissociation stimulator           949 TAATTTGCGTT         4,00         1,56 Hs.79368 epithelial membrane protein 1           950 AGAATCGTTTG         4,00         1,56 Hs.136299 ESTs           951 AACAGTCAAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.998135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.99858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 Procollagen C-endopeptidase enhance
947 AGGGGAAGGTG         8,00         2,92 Hs.112540 EST           948 CAAGACGGGG         19,00         6,29 Hs.106185 ral guanine nucleotide dissociation stimulator           949 TAATTTGCGTT         4,00         1,56 Hs.79368 epithelial membrane protein 1           950 AGAATCGTTTG         4,00         1,56 Hs.136299 ESTs           951 AACAGTCAAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALUB_HUMAN ALU SUBFAMILY           953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.99858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 procollagen C-endopeptidase enhancer
948         CAAGACGGGG         19,00         6,29         Hs.106185         ral guanine nucleotide dissociation stimulator           949         TAATTTGCGTT         4,00         1,56         Hs.79368         epithelial membrane protein 1           950         AGAATCGTTTG         4,00         1,56         Hs.136299         ESTs           951         AACAGTCAAAA         17,00         5,66         Hs.26557         plakophilin 3           952         GCAAAACTCTG         10,00         3,49         Hs.278746         ESTs, Moderately similar to ALUB_HUMAN ALU SUBFAMILY           953         GCCAGCCAGTG         10,00         3,49         Hs.149098         smoothelin           954         GCGAAATCCCA         6,00         2,21         Hs.268728         ESTs           955         TCTGTGGTCCC         6,00         2,21         Hs.27894         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956         CCTATAATTCC         6,00         2,21         Hs.135491         ESTs           957         GCAATAAAATA         1,00         0,40         Hs.98135         ESTs           958         TGGAATCCAGG         1,00         0,40         Hs.89474         ADP-ribosylation factor 6           961         ATGGAGGTATG
Stimulator   949 TAATTTGCGTT   4,00   1,56 Hs.79368   epithelial membrane protein 1   950 AGAATCGTTTG   4,00   1,56 Hs.136299   ESTs   951 AACAGTCAAAA   17,00   5,66 Hs.26557   plakophilin 3   952 GCAAAACTCTG   10,00   3,49 Hs.278746   ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY   953 GCCAGCCAGTG   10,00   3,49 Hs.149098   smoothelin   954 GCGAAATCCCA   6,00   2,21 Hs.268728   ESTs   955 TCTGTGGTCCC   6,00   2,21 Hs.227894   ESTs, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J S   956 CCTATAATTCC   6,00   2,21 Hs.135491   ESTs   957 GCAATAAAATA   1,00   0,40 Hs.99621   ESTs   958 TGGAATCCAGG   1,00   0,40 Hs.98135   ESTs   959 CTTTTGTCAGC   1,00   0,40 Hs.9858   Homo sapiens clone 25023 mRNA   sequence   960 TTATTGTATTG   1,00   0,40 Hs.89444   ADP-ribosylation factor 6   961 ATGGAGGTATG   1,00   0,40 Hs.89444   procollagen C-endopeptidase enhancer   2
949 TAATTTGCGTT         4,00         1,56 Hs.79368         epithelial membrane protein 1           950 AGAATCGTTTG         4,00         1,56 Hs.136299 ESTs           951 AACAGTCAAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 procollagen C-endopeptidase enhancer
950 AGAATCGTTTG         4,00         1,56 Hs.136299 ESTs           951 AACAGTCAAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 procollagen C-endopeptidase enhancer
951 AACAGTCAAAA         17,00         5,66 Hs.26557 plakophilin 3           952 GCAAAACTCTG         10,00         3,49 Hs.278746 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 procollagen C-endopeptidase enhancer
952         GCAAAACTCTG         10,00         3,49         Hs.278746         ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           953         GCCAGCCAGTG         10,00         3,49         Hs.149098         smoothelin           954         GCGAAATCCCA         6,00         2,21         Hs.268728         ESTs           955         TCTGTGGTCCC         6,00         2,21         Hs.227894         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956         CCTATAATTCC         6,00         2,21         Hs.135491         ESTs           957         GCAATAAAATA         1,00         0,40         Hs.99621         ESTs           958         TGGAATCCAGG         1,00         0,40         Hs.998135         ESTs           959         CTTTTGTCAGC         1,00         0,40         Hs.90858         Homo sapiens clone 25023 mRNA sequence           960         TTATTGTATTG         1,00         0,40         Hs.89474         ADP-ribosylation factor 6           961         ATGGAGGTATG         1,00         0,40         Hs.89444         procollagen C-endopeptidase enhancer           2
ALU8_HUMAN ALU SUBFAMILY
953 GCCAGCCAGTG         10,00         3,49 Hs.149098 smoothelin           954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 procollagen C-endopeptidase enhancer
954 GCGAAATCCCA         6,00         2,21 Hs.268728 ESTs           955 TCTGTGGTCCC         6,00         2,21 Hs.227894 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           956 CCTATAATTCC         6,00         2,21 Hs.135491 ESTs           957 GCAATAAAATA         1,00         0,40 Hs.99621 ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135 ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858 Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474 ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944 procollagen C-endopeptidase enhancer
ALU SUBFAMILY J S
956         CCTATAATTCC         6,00         2,21         Hs.135491         ESTs           957         GCAATAAAATA         1,00         0,40         Hs.99621         ESTs           958         TGGAATCCAGG         1,00         0,40         Hs.98135         ESTs           959         CTTTTGTCAGC         1,00         0,40         Hs.90858         Homo sapiens clone 25023 mRNA sequence           960         TTATTGTATTG         1,00         0,40         Hs.89474         ADP-ribosylation factor 6           961         ATGGAGGTATG         1,00         0,40         Hs.8944         procollagen C-endopeptidase enhancer
957 GCAATAAAATA         1,00         0,40 Hs.99621         ESTs           958 TGGAATCCAGG         1,00         0,40 Hs.98135         ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858         Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474         ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944         procollagen C-endopeptidase enhancer
958 TGGAATCCAGG         1,00         0,40 Hs.98135         ESTs           959 CTTTTGTCAGC         1,00         0,40 Hs.90858         Homo sapiens clone 25023 mRNA sequence           960 TTATTGTATTG         1,00         0,40 Hs.89474         ADP-ribosylation factor 6           961 ATGGAGGTATG         1,00         0,40 Hs.8944         procollagen C-endopeptidase enhancer 2
959 CTTTTGTCAGC 1,00 0,40 Hs.90858 Homo sapiens clone 25023 mRNA sequence 960 TTATTGTATTG 1,00 0,40 Hs.89474 ADP-ribosylation factor 6 961 ATGGAGGTATG 1,00 0,40 Hs.8944 procollagen C-endopeptidase enhancer 2
Sequence   Sequence
960 TTATTGTATTG 1,00 0,40 Hs.89474 ADP-ribosylation factor 6 961 ATGGAGGTATG 1,00 0,40 Hs.8944 procollagen C-endopeptidase enhancer 2
961 ATGGAGGTATG 1,00 0,40 Hs.8944 procollagen C-endopeptidase enhancer 2
2
962 TCGTGTTTTCG   1,00  0,40 Hs.87595  translocase of inner mitochondrial
membrane 22 (yeas
963 TTGTGGCCCCA 1,00 0,40 Hs.84630 ESTs
964 TTTGTTTGTTT 1,00 0,40 Hs.8355 ESTs
965 GTCACGAACAT 1,00 0,40 Hs.82933 protein x 013
966 GTTAAGGTAAA 1,00 0,40 Hs.79241 B-cell CLL/lymphoma 2
967 AAGAAGAAAG 1,00 0,40 Hs.78293 ESTs
968 TAGCCAGTTAA 1,00 0,40 Hs.74101 spleen tyrosine kinase
969 CTGCCGGGGC 1,00 0,40 Hs.74097 mercaptopyruvate sulfurtransferase
970 AACGGTGTTTG 1,00 0,40 Hs.71371 ESTs
971 CATAAACGGC 1,00 0,40 Hs.69954   laminin, gamma 3

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973 CTGGCATAGAA 1,00 0,4 974 ACTGCCCTGA 1,00 0,4	0 Hs.69743 0 Hs.61272	GM2 ganglioside activator protein
974 ACTGCCCCTGA 1,00 0,4		IECT-
<del></del>		ESTs
975 TCTGCTGCCTG   1,00 0,4	<del></del>	semaphorin sem2 ESTs
<del></del>		
	0 Hs.54982	ESTs, Weakly similar to ALU4_HUMAN ALU SUBFAMILY SB2
		ESTs
		DKFZP434B168 protein
	0 Hs.47587	ESTs
		ESTs
		leukotriene C4 synthase
		ESTs
		ESTs
984 ACAACTGGAAT   1,00   0,4		Homo sapiens DNA binding peptide mRNA, partial cds
985 TAATAAAATGC 1,00 0,4		ESTs
	Hs.284153	Fanconi anemia, complementation group A
987 TTTTGAAGATA 1,00 0,4	Hs.283322	hypothetical protein
988 GCAAATCAGAT 1,00 0,4	Hs.279477	ESTs
	Hs.277896	
	Hs.276994	
	Hs.270331	
	Hs.269838	
		natural killer-tumor recognition sequence
		chondroitin 4-sulfotransferase
	Hs.236505	EST, Weakly similar to ALU6_HUMAN ALU SUBFAMILY SP S
996 GTTTATAATTA 1,00 0,4	Hs.231966	ESTs
997 CTGGCACCCTG 1,00 0,4	Hs.212716	EST
	Hs.204930	
999 GAGCAGGCAAA 1,00 0,4	Hs.200333	apolipoprotein B48 receptor
1000 CCAAAAAGTG 1,00 0,4		sterol-C5-desaturase (fungal ERG3, delta-5-desaturas
1001 CCAGAGGAATG 1,00 0,4	Hs.180414	heat shock 70kd protein 10 (HSC71)
	Hs.179091	
	Hs.174031	cytochrome c oxidase subunit VIb
	Hs.17258	
	Hs.16959	ESTs
	Hs.166293	
1007 TATATAAGTAC 1,00 0,4		catenin (cadherin-associated protein), delta 1
1008 TAATAATACAA 1,00 0,4		KIAA0431 protein
		hypothetical protein FLJ20159
		alpha-2-plasmin inhibitor
		Homo sapiens clone 23728 mRNA
1 1,33		sequence
1012 CTGTTTTGAA 1,00 0,4		M-phase phosphoprotein 6
		DKFZP727G051 protein

1015   CCTACAGTCCC						
DKFZp761C0524 (from clone DK			1,00			
1016   CAATCCTCCTG	1015	CCTACAGTCCC	1,00	0,40		
1017   AACAAGTAATA						
1018   TCCAGTACAGA						
ID21   TITCTGRAGTG			1,00			
1022   TCCTGTGATTT	1020	ACTCTGCTCGG				
1023   AAAGCACAAGT	1021	TTCTGTGAGTG	1,00	0,40	Hs.122559	ESTs
1024   TTTTGAAAAA	1022	TCCTGTGATTT	1,00	0,40	Hs.12253	ESTs
Deta subcomplex, 6     Deta subcomplex, 6	1023	AAAGCACAAGT	1,00	0,40	Hs.111758	keratin 6B
1025   AAGTCCTGGCC	1024	TTTTGAAAAA	1,00	0,40	Hs.109646	NADH dehydrogenase (ubiquinone) 1
Motif-binding fa   Motif-binding fa   ESTs, Weakly similar to spastin protein   H.sapiens   ESTs, Weakly similar to spastin protein   H.sapiens   ESTs, Meakly similar to spastin protein   H.sapiens   ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY   Moderately similar to ALU1_HUMAN ALU SUBFAMILY   Motifical protein PRO2859   Motifical protein protei			1			beta subcomplex, 6
Motif-binding fa   Motif-binding fa   ESTs, Weakly similar to spastin protein   H.sapiens   ESTs, Weakly similar to spastin protein   H.sapiens   ESTs, Meakly similar to spastin protein   H.sapiens   ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY   Moderately similar to ALU1_HUMAN ALU SUBFAMILY   Motifical protein PRO2859   Motifical protein protei	1025	AAGTCCTGGCC	1,00	0,40	Hs.109314	ESTs, Weakly similar to 2202255A AT
[H.sapiens]						motif-binding fa
[H.sapiens]	1026	TGTTACTGGAT	1,00	0,40	Hs.100861	ESTs, Weakly similar to spastin protein
ALU1_HUMAN ALU SUBFAMILY						
1028   GTGGCGGCAC   85,00   25,22   Hs.283044   hypothetical protein PRO2859   1029   CCACTTGCACT   3,00   1,18   Hs.220962   EST, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J SE   1030   TTCCATACCCC   3,00   1,18   Hs.180398   LIM domain-containing preferred translocation partne   1031   ATTGCATCACT   5,00   1,85   Hs.209111   EST   major histocompatibility complex, class   II, DR   beta   II, DR   beta   III, DR   beta	1027	CCTGTAATCTT	8,00	2,84	Hs.120882	ESTs, Moderately similar to
1029   CCACTTGCACT   3,00						ALU1_HUMAN ALU SUBFAMILY
1029   CCACTTGCACT   3,00	1028	GTGGCGGGCAC	85,00	25,22	Hs.283044	hypothetical protein PRO2859
1,18	1029	CCACTTGCACT	3,00	1,18	Hs.220962	EST, Weakly similar to ALU1_HUMAN
translocation partne   translocation partner   translocation partner   translocation partner   translocation partner   translocation partner   translocation partner   translocation						ALU SUBFAMILY J SE
1,85   Hs.209111   EST   major histocompatibility complex, class   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   ESTs, Weakly similar to Pl-3 kinase   II, DR beta   II, D	1030	TTCCATACCCC	3,00	1,18	Hs.180398	LIM domain-containing preferred
1032 GAGTTAAAAAA						translocation partne
II, DR beta   II, DR beta   II, DR beta   III, DR beta   IIII, DR beta   IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	1031	ATTGCATCACT	5,00			
1,53	1032	GAGTTAAAAAA	5,00	1,85	Hs.180255	major histocompatibility complex, class
[H.sapiens]  1034 CAGATGGAGGC						
1034   CAGATGGAGGC	1033	GGGCCCTGGCC	4,00	1,53	Hs.25895	ESTs, Weakly similar to PI-3 kinase
1035   CCTCTCCCACA   2,00   0,82   Hs.99197   ESTs     1036   GCCAGGGGGTA   2,00   0,82   Hs.96875   ESTs     1037   CTCAGTCTTTT   2,00   0,82   Hs.76722   CCAAT/enhancer binding protein (C/EBP), delta     1038   GATGTATTCTA   2,00   0,82   Hs.75844   ESTs, Highly similar to AF151903_1     1039   CCCTTCTGTAA   2,00   0,82   Hs.75716   plasminogen activator inhibitor, type II (arginine-s     1040   TGGAACTGTCA   2,00   0,82   Hs.285802   ESTs     1041   GGCCTGTAATC   2,00   0,82   Hs.267400   ESTs, Moderately similar to ALUT_HUMAN ALU SUBFAMILY     1042   AGGTATATATC   2,00   0,82   Hs.24715   Homo sapiens mRNA; cDNA     1043   TTCTGAAAGGA   2,00   0,82   Hs.227209   DKFZP586F1019 protein     1044   AGCCTGTGGTC   2,00   0,82   Hs.154919   KIAA0625 protein     1045   TTGCGTGTGTC   2,00   0,82   Hs.1183   dual specificity phosphatase 2     1046   CATAATTTCTC   2,00   0,82   Hs.104660   EF-5A2 protein     1047   TGGGACGTGAG   7,00   2,45   Hs.3796   EphB6						[H.sapiens]
1035   CCTCTCCCACA   2,00   0,82   Hs.99197   ESTs     1036   GCCAGGGGGTA   2,00   0,82   Hs.96875   ESTs     1037   CTCAGTCTTTT   2,00   0,82   Hs.76722   CCAAT/enhancer binding protein (C/EBP), delta     1038   GATGTATTCTA   2,00   0,82   Hs.75844   ESTs, Highly similar to AF151903_1     1039   CCCTTCTGTAA   2,00   0,82   Hs.75716   plasminogen activator inhibitor, type II (arginine-s     1040   TGGAACTGTCA   2,00   0,82   Hs.285802   ESTs     1041   GGCCTGTAATC   2,00   0,82   Hs.267400   ESTs, Moderately similar to ALUT_HUMAN ALU SUBFAMILY     1042   AGGTATATATC   2,00   0,82   Hs.24715   Homo sapiens mRNA; cDNA     1043   TTCTGAAAGGA   2,00   0,82   Hs.227209   DKFZP586F1019 protein     1044   AGCCTGTGGTC   2,00   0,82   Hs.154919   KIAA0625 protein     1045   TTGCGTGTGTC   2,00   0,82   Hs.1183   dual specificity phosphatase 2     1046   CATAATTTCTC   2,00   0,82   Hs.104660   EF-5A2 protein     1047   TGGGACGTGAG   7,00   2,45   Hs.3796   EphB6	1034	CAGATGGAGGC	4,00	1,53	Hs.127273	hypothetical protein FLJ10044
1037   CTCAGTCTTTT   2,00   0,82   Hs.76722   CCAAT/enhancer binding protein (C/EBP), delta   1038   GATGTATTCTA   2,00   0,82   Hs.75844   ESTs, Highly similar to AF151903_1   CGI-145 protein [   plasminogen activator inhibitor, type II (arginine-s   1040   TGGAACTGTCA   2,00   0,82   Hs.285802   ESTs   1041   GGCCTGTAATC   2,00   0,82   Hs.267400   ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY   1042   AGGTATATATC   2,00   0,82   Hs.24715   Homo sapiens mRNA; cDNA DKFZP434D0215 (from clone DK   1043   TTCTGAAAGGA   2,00   0,82   Hs.227209   DKFZP586F1019 protein   1044   AGCCTGTGGTC   2,00   0,82   Hs.154919   KIAA0625 protein   1045   TTGCGTGTGTC   2,00   0,82   Hs.1183   dual specificity phosphatase 2   1046   CATAATTTCTC   2,00   0,82   Hs.104660   eIF-5A2 protein   1047   TGGGACGTGAG   7,00   2,45   Hs.3796   EphB6	1035	CCTCTCCCACA	2,00	0,82	Hs.99197	ESTs
C/EBP), delta   C/EBP), delt	1036	GCCAGGGGGTA	2,00	0,82	Hs.96875	ESTs
1038 GATGTATTCTA	1037	CTCAGTCTTTT	2,00	0,82	Hs.76722	CCAAT/enhancer binding protein
CGI-145 protein						(C/EBP), delta
1039   CCCTTCTGTAA   2,00   0,82   Hs.75716   plasminogen activator inhibitor, type II (arginine-s   1040   TGGAACTGTCA   2,00   0,82   Hs.285802   ESTs   1041   GGCCTGTAATC   2,00   0,82   Hs.267400   ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY   1042   AGGTATATATC   2,00   0,82   Hs.24715   Homo sapiens mRNA; cDNA DKFZp434D0215 (from clone DK   1043   TTCTGAAAGGA   2,00   0,82   Hs.227209   DKFZP586F1019 protein   1044   AGCCTGTGGTC   2,00   0,82   Hs.154919   KIAA0625 protein   1045   TTGCGTGTGTC   2,00   0,82   Hs.1183   dual specificity phosphatase 2   1046   CATAATTTCTC   2,00   0,82   Hs.104660   eIF-5A2 protein   1047   TGGGACGTGAG   7,00   2,45   Hs.3796   EphB6	1038	GATGTATTCTA	2,00	0,82	Hs.75844	ESTs, Highly similar to AF151903_1
(arginine-s						CGI-145 protein [
(arginine-s	1039	CCCTTCTGTAA	2,00	0,82	Hs.75716	plasminogen activator inhibitor, type II
1041   GGCCTGTAATC   2,00   0,82   Hs.267400   ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY     1042   AGGTATATATC   2,00   0,82   Hs.24715   Homo sapiens mRNA; cDNA DKFZp434D0215 (from clone DK     1043   TTCTGAAAGGA   2,00   0,82   Hs.227209   DKFZP586F1019 protein     1044   AGCCTGTGGTC   2,00   0,82   Hs.154919   KIAA0625 protein     1045   TTGCGTGTGTC   2,00   0,82   Hs.1183   dual specificity phosphatase 2     1046   CATAATTTCTC   2,00   0,82   Hs.104660   eIF-5A2 protein     1047   TGGGACGTGAG   7,00   2,45   Hs.3796   EphB6						
ALU7_HUMAN ALU SUBFAMILY	1040	TGGAACTGTCA	2,00	0,82	Hs.285802	ESTs
ALU7_HUMAN ALU SUBFAMILY	1041	GGCCTGTAATC	2,00	0,82	Hs.267400	ESTs, Moderately similar to
DKFZp434D0215 (from clone DK						
1043 TTCTGAAAGGA         2,00         0,82 Hs.227209 DKFZP586F1019 protein           1044 AGCCTGTGGTC         2,00         0,82 Hs.154919 KIAA0625 protein           1045 TTGCGTGTGTC         2,00         0,82 Hs.1183 dual specificity phosphatase 2           1046 CATAATTTCTC         2,00         0,82 Hs.104660 eIF-5A2 protein           1047 TGGGACGTGAG         7,00         2,45 Hs.3796 EphB6	1042	AGGTATATATC	2,00	0,82	Hs.24715	Homo sapiens mRNA; cDNA
1044 AGCCTGTGGTC         2,00         0,82 Hs.154919 KIAA0625 protein           1045 TTGCGTGTC         2,00         0,82 Hs.1183 dual specificity phosphatase 2           1046 CATAATTTCTC         2,00         0,82 Hs.104660 eIF-5A2 protein           1047 TGGGACGTGAG         7,00         2,45 Hs.3796 EphB6			3			DKFZp434D0215 (from clone DK
1045 TTGCGTGTC         2,00         0,82 Hs.1183         dual specificity phosphatase 2           1046 CATAATTTCTC         2,00         0,82 Hs.104660 eIF-5A2 protein           1047 TGGGACGTGAG         7,00         2,45 Hs.3796         EphB6	1043	TTCTGAAAGGA	2,00	0,82	Hs.227209	
1045 TTGCGTGTC         2,00         0,82 Hs.1183         dual specificity phosphatase 2           1046 CATAATTTCTC         2,00         0,82 Hs.104660 eIF-5A2 protein           1047 TGGGACGTGAG         7,00         2,45 Hs.3796         EphB6	1044	AGCCTGTGGTC	2,00	0,82	Hs.154919	KIAA0625 protein
1046 CATAATTTCTC 2,00 0,82 Hs.104660 eIF-5A2 protein 1047 TGGGACGTGAG 7,00 2,45 Hs.3796 EphB6	1045	TTGCGTGTGTC				
1047 TGGGACGTGAG 7,00 2,45 Hs.3796 EphB6						
107010001 V 110000 1 10,001 0,071110,10720 HEG 1a. VICARIV SIIIIIIGI IO AEGE 1101111 II V		GCGAAATCCCG	10,00			<del></del>

		<del></del>		···-	
					ALU SUBFAMILY SB
1049	ACCAAAAACCA	120,00			collagen, type I, alpha 1
1050	AACCCAGGAGG	126,00			hypothetical protein FLJ20159
	TCTCTGTGTAG	4,00	1,49	Hs.79187	coxsackie virus and adenovirus receptor
1052	CGCAGTAGGGG	4,00		Hs.17411	KIAA0699 protein
1053	CGAGAGGGAGA	4,00	1,49	Hs.158159	FAT tumor suppressor (Drosophila)
1071	0-0001-0-00		4.40	11 (011	homolog 2
	GTGGCATCTGC	4,00		Hs.1244	CD9 antigen (p24)
1055	CTAACGGGGCG	7,00	2,42	Hs.102171	immunoglobulin superfamily containing leucine-rich r
1056	GCAAAACCCCT	6,00	2,10	Hs.75238	chromatin assembly factor 1, subunit B (p60)
1057	ACTGCTTTACT	6,00	2.10	Hs.72157	DKFZP564I1922 protein
	CCCCAGGCTGC	3,00		Hs.9645	ESTs
	TAAAATGTTTA	3,00			ESTs
	CCTACTGCACT	3,00			ESTs, Moderately similar to KIAA0680 protein [H.sapi
1061	GTGGCTCATTC	3,00	1.15	Hs.116577	prostate differentiation factor
	AAGCACAAAAA	12,00		Hs.9963	TYRO protein tyrosine kinase binding protein
1063	GTGGCGGGCGC	94,00	26 43	Hs 129710	malignancy-associated protein
	AGAACCTTAAA	14,00			major histocompatibility complex, class I,
1065	CCTGAAATCCC	8,00	2.60	Hs.182124	EST <sub>2</sub>
	GCGAAACCCAG				
	ATGTAGGTGCC	13,00			HP1-BP74
		9,00			phosphatidic acid phosphatase type 2B
	TTAAATAGCAC TTTTATTTCCA	10,00		Hs. 172920	collagen, type I, alpha 1
		4,00			ESTs, Weakly similar to ALUB_HUMAN IIII ALU CLASS B
1070	AATTACAGCCA	4,00	5,89	Hs.74471	gap junction protein, alpha 1, 43kD (connexin 43)
1071	CCTTACCTAAG	4,00	5,89	Hs.240217	dopachrome tautomerase (dopachrome delta-isomerase,
1072	CTCCCTGAACG	4,00	5.89	Hs.11006	ESTs
	AACACGAATGA	4,00		Hs.259855	
	GTGGCAAGCAC	4,00			Rho GTPase activating protein 1
	TTCACCATCCT	4,00			ESTs, Weakly similar to tetraspan NET-4 [H.sapiens]
1076	AGAGGGAGTGA	2,00	0,78	Hs.85201	C-type (calcium dependent, carbohydrate-recognition
1077	AGTCCTTGAAA	2,00	0,78	Hs.81665	v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogen
1078	CCCAGCCTAAA	2,00	0,78	Hs.47986	Homo sapiens mRNA; cDNA DKFZp586H051 (from clone DKF
1079	TTTAACTGACA	2,00	0.78	Hs.24880	ESTs
	CCTTGTAATCC	2,00		Hs.197054	
	GAAGGCTTATC	2,00			nuclear factor of activated T-cells,
1082	TAGCAAAGATT	2.00	0.79	He 166172	cytoplasmic 3 aryl hydrocarbon receptor nuclear
1002	INGONAGAII	2,00	0,70	<u>   13. 100 1/2</u>	aryr nyurocarbon receptor nuclear

					ltennologica
4000	CACATOTOTAA	0.00	0.70	115 40500	translocator
	CACATCTGTAA	2,00			myosin phosphatase, target subunit 1
	TTCAGTAATAA	2,00			hypothetical protein FLJ20847
	GGTGAAACCCT	2,00			CGI-43 protein
	CTTCTGCCTCA	2,00		Hs.115896	
1087	GAGAGGTGATT	2,00	0,78	Hs.114062	protein tyrosine phosphatase-like (proline
					instead o
	ATGTATTTTA	2,00			ALR-like protein
1089	CTATAGGAGAC	5,00			integral membrane protein 1
1090	TCCGTGTATAA	5,00	1,75	Hs.3321	ESTs, Highly similar to IRX3_MOUSE
					IROQUOIS-CLASS HO
1091	GCAAAACCCCA	98,00	26,53		tumor necrosis factor (ligand)
					superfamily, member 1
1092	TTCCATAGCCT	6,00			Notch (Drosophila) homolog 3
	CTGTGAAATGC	3,00			hypothetical protein FLJ10704
1094	AAAGAACATAG	3,00	<u> </u>	Hs.104558	ESTs
1095	CACCTGTAATC	37,00		Hs.275819	
1096	GGCAACAAGAG	9,00	2,89	Hs.205739	ESTs, Weakly similar to ALU7_HUMAN
					ALU SUBFAMILY SQ
1097	GTGGCGGGTGC	72,00	19,32	Hs.277015	EST
1098	ACCTTCAAAAA	4,00		Hs.28444	hypothetical protein FLJ10567
1099	ACATCTGGCTT	4,00			KIAA0737 gene product
-	GTACGTATTCT	6,00			immunoglobulin J polypeptide, linker
			·		protein for imm
1101	ATCCGCCTGCC	6,00	2,00	Hs. 167956	ESTs, Weakly similar to KIAA0309
		, -			[H.sapiens]
1102	ACCCACGTCAG	24,00	6,84		jun B proto-oncogene
	TAATCCCAGCT	14,00		Hs.238384	
	ATTGCACCACT	46,00			CGI-43 protein
	AGGACCAAGGA	1,00			ESTs, Weakly similar to ALU7_HUMAN
		.,	•,•		ALU SUBFAMILY SQ
1106	TAAGCTACTAA	1,00	0.37	Hs.97469	ESTs, Weakly similar to I49698 alpha-
		.,	-,		1,3-galactosylt
1107	CTCCATTGTCT	1,00	0.37	Hs.93005	slug (chicken homolog), zinc finger
		.,	-,-:		protein
1108	TGACATTAAAC	1,00	0.37	Hs.87432	ESTs
	GGATTCAAGAG	1,00		Hs.86947	a disintegrin and metalloproteinase
' ' '		.,	٠,٠.		domain 8
1110	ATGTTATCATA	1,00	0.37	Hs.8325	mitogen-activated protein kinase 9
	TACTCTGTTGA	1,00		Hs.82587	phospholipase D1, phophatidylcholine-
ļ		.,55	3,01		specific
1112	GGAAAAGAAAA	1,00	0.37	Hs.82141	Human clone 23612 mRNA sequence
	TATACGTTATG	1,00		Hs.78894	KIAA0161 gene product
	CACTTGGTGAT	1,00		Hs.77348	hydroxyprostaglandin dehydrogenase
1'''	S.O. TOOTOAT	',55	0,57	1.3.7 7 3-40	15-(NAD)
1115	ATGAATGTAAA	1,00	0.37	Hs.76853	Homo sapiens mRNA full length insert
' ' ' '		',00	0,37	1 13.7 0000	cDNA clone EURO
1116	TGTTTTCATAA	1,00	0.37	Hs.75703	small inducible cytokine A4 (homologous
1,,,,		',00	0,57	1 13.7 37 03	to mouse Mip
		L		L	Iro monde mih

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1118   GTGGTTCATTC				<del> </del>		
Sulfotransfer   Sulfotransfer   Sing   Sulfotransfer   Sing   S			1,00			cytochrome c oxidase subunit VIc
1119   CAACAAAAGCA	1118	GTGGTTCATTC	1,00	0,37	Hs.6853	
1120   CATTITITIGG	1119	CAACAAAAGCA	1,00	0,37	Hs.66450	
1121   CTGCTAAACTA	1120	CATTTTTGCG				ESTs
1122   ACCCTGAATGG				0.37	Hs.46826	ESTs
1123   TTGTAACAAAA						
1124   TGCTGTTCATA						jumonji (mouse) homolog
1125   TATGTGGGTTA	_					
1126   CAGCAATTATA	1125	TATGTGGGTTA		0,37	Hs.34359	ESTs
1127   CTCCATTGCCA	1126	CAGCAATTATA		0,37	Hs.32309	inositol polyphosphate-1-phosphatase
1129   ATTCCACCACT	1127	CTCCATTGCCA	1,00	0,37	Hs.31869	
1130   TCTTACTCAGA   1,00   0,37   Hs.285081   ESTs   Homo sapiens PAK2 mRNA, complete cds   1131   AATAAAAAATA   1,00   0,37   Hs.284275   Homo sapiens PAK2 mRNA, complete cds   1132   GAGACAGTGAC   1,00   0,37   Hs.284146   hypothetical protein DKFZp762N0610   1133   CCTGGGCAACA   1,00   0,37   Hs.273683   EST   1134   CCTTTGCACTC   1,00   0,37   Hs.265124   ESTs   1135   TGCAGACAGGG   1,00   0,37   Hs.264363   hypothetical protein FLJ10110   Rho guanine nucleotide exchange facto (GEF) 3   1137   TCTTTAAAGTA   1,00   0,37   Hs.25951   guanine nucleotide regulatory protein (oncogene)   1138   ATATTGGTGGT   1,00   0,37   Hs.250692   hepatic leukemia factor   1139   GTGGCGAATGC   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1141   TCTCCATTCCT   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1142   TATATAGAATG   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1142   TATATAGAATG   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1144   CACTCCAACCT   1,00   0,37   Hs.22581   ESTs   ESTs   Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S   1144   CACTCCAACCT   1,00   0,37   Hs.205899   ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC   myosin regulatory light chain interacting protein   1146   CCTTCTGAATA   1,00   0,37   Hs.194660   Cercid-lipofuscinosis, neuronal 3, juvenii   1148   ATTGTACAACA   1,00   0,37   Hs.194660   Cercid-lipofuscinosis, neuronal 3, juvenii   1148   ATTGTACAACA   1,00   0,37   Hs.184326   CDC10 (cell division cycle 10, S. cerevisiae, homolo   1149   AGCCTATTAAA   1,00   0,37   Hs.183902   ESTs   1150   GCCCCTGCGCC   1,00   0,37   Hs.183002   ESTs   1150   GCCCCTGCGCC   1,00   0,37   Hs.176065   ESTs   1152   TTCATTAAGAA   1,00   0,37   Hs.17411   KIAA0699 protein   1150   TCCATTAAA   1,00   0,37   Hs.17411   KIAA0699 protein   1150   TCCATTAAA   1,00   0,37   Hs.17411   TCCATTAACAA   1,00   0,37   Hs.17411   TCCATTAACAA   1,00   0,37   Hs.17411   TCCATTAAC	1128	GGGTGGGTCAC	1,00	0,37	Hs.31500	ESTs
1130   TCTTACTCAGA   1,00   0,37   Hs.285081   ESTs   Homo sapiens PAK2 mRNA, complete cds   1131   AATAAAAAATA   1,00   0,37   Hs.284275   Homo sapiens PAK2 mRNA, complete cds   1132   GAGACAGTGAC   1,00   0,37   Hs.284146   hypothetical protein DKFZp762N0610   1133   CCTGGGCAACA   1,00   0,37   Hs.273683   EST   1134   CCTTTGCACTC   1,00   0,37   Hs.265124   ESTs   1135   TGCAGACAGGG   1,00   0,37   Hs.264363   hypothetical protein FLJ10110   Rho guanine nucleotide exchange facto (GEF) 3   1137   TCTTTAAAGTA   1,00   0,37   Hs.25951   guanine nucleotide regulatory protein (oncogene)   1138   ATATTGGTGGT   1,00   0,37   Hs.250692   hepatic leukemia factor   1139   GTGGCGAATGC   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1141   TCTCCATTCCT   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1142   TATATAGAATG   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1142   TATATAGAATG   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i   1144   CACTCCAACCT   1,00   0,37   Hs.22581   ESTs   ESTs   Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S   1144   CACTCCAACCT   1,00   0,37   Hs.205899   ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC   myosin regulatory light chain interacting protein   1146   CCTTCTGAATA   1,00   0,37   Hs.194660   Cercid-lipofuscinosis, neuronal 3, juvenii   1148   ATTGTACAACA   1,00   0,37   Hs.194660   Cercid-lipofuscinosis, neuronal 3, juvenii   1148   ATTGTACAACA   1,00   0,37   Hs.184326   CDC10 (cell division cycle 10, S. cerevisiae, homolo   1149   AGCCTATTAAA   1,00   0,37   Hs.183902   ESTs   1150   GCCCCTGCGCC   1,00   0,37   Hs.183002   ESTs   1150   GCCCCTGCGCC   1,00   0,37   Hs.176065   ESTs   1152   TTCATTAAGAA   1,00   0,37   Hs.17411   KIAA0699 protein   1150   TCCATTAAA   1,00   0,37   Hs.17411   KIAA0699 protein   1150   TCCATTAAA   1,00   0,37   Hs.17411   TCCATTAACAA   1,00   0,37   Hs.17411   TCCATTAACAA   1,00   0,37   Hs.17411   TCCATTAAC	1129	ATTCCACCACT	1,00	0,37	Hs.2934	ribonucleotide reductase M1 polypeptide
1132   GAGACAGTGAC	1130	TCTTACTCAGA	1,00	0,37		
1133   CCTGGGCAACA   1,00   0,37   Hs.265124   ESTs     1134   CCTTTGCACTC   1,00   0,37   Hs.264363   hypothetical protein FLJ10110     1136   TTGGCCCAGTC   1,00   0,37   Hs.25951   Rho guanine nucleotide exchange facto (GEF) 3     1137   TCTTTAAAGTA   1,00   0,37   Hs.25155   guanine nucleotide regulatory protein (oncogene)     1138   ATATTGGTGGT   1,00   0,37   Hs.250692   hepatic leukemia factor     1139   GTGGCGAATGC   1,00   0,37   Hs.230479   EST     1140   TATTAACATTC   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i     1141   TCTCCATTCCT   1,00   0,37   Hs.226573   inhibitor of kappa light polypeptide gene enhancer i     1142   TATATAGAATG   1,00   0,37   Hs.22581   ESTs   ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S     1144   CACTCCAACCT   1,00   0,37   Hs.205899   ESTS, Weakly similar to ALU5_HUMAN ALU SUBFAMILY J S     1145   CGATGTTAAAA   1,00   0,37   Hs.205899   ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC   myosin regulatory light chain interacting protein     1147   TGAGGATCCAG   1,00   0,37   Hs.194660   Ceroid-lipofuscinosis, neuronal 3, juvenil (Batten,     1148   ATTGTACAACA   1,00   0,37   Hs.184326   CDC10 (cell division cycle 10, S. cerevisiae, homolo     1149   AGCCTATTAAA   1,00   0,37   Hs.183593   zinc finger protein 24 (KOX 17)     1151   GTGAATGCACT   1,00   0,37   Hs.183092   ESTs     1151   GTGAATGCACT   1,00   0,37   Hs.18065   ESTs     1152   TTCATTAAGAA   1,00   0,37   Hs.17411   KIAA0699 protein	1131	AATAAAAAATA	1,00	0,37	Hs.284275	
1134   CCTTTGCACTC	1132	GAGACAGTGAC	1,00	0,37	Hs.284146	hypothetical protein DKFZp762N0610
1135   TGCAGACAGGG	1133	CCTGGGCAACA	1,00	0,37	Hs.273683	EST
1136 TTGGCCCAGTC	1134	CCTTTGCACTC	1,00	0,37	Hs.265124	ESTs
CGEF) 3			1,00	0,37		
(oncogene)	1136	TTGGCCCAGTC	1,00	0,37	Hs.25951	
1139 GTGGCGAATGC         1,00         0,37 Hs.230479 EST           1140 TATTAACATTC         1,00         0,37 Hs.226573 inhibitor of kappa light polypeptide gene enhancer i           1141 TCTCCATTCCT         1,00         0,37 Hs.226573 inhibitor of kappa light polypeptide gene enhancer i           1142 TATATAGAATG         1,00         0,37 Hs.22581 ESTs           1143 GCGAGATCCTG         1,00         0,37 Hs.22529 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1144 CACTCCAACCT         1,00         0,37 Hs.205899 ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC           1145 CGATGTTAAAA         1,00         0,37 Hs.20072 myosin regulatory light chain interacting protein           1146 CCTTCTGAATA         1,00         0,37 Hs.194660 ceroid-lipofuscinosis, neuronal 3, juvenil (Batten,           1147 TGAGGATCCAG         1,00         0,37 Hs.191621 ESTs           1148 ATTGTACAACA         1,00         0,37 Hs.184326 CDC10 (cell division cycle 10, S. cerevisiae, homolo           1149 AGCCTATTAAA         1,00         0,37 Hs.183593 zinc finger protein 24 (KOX 17)           1150 GCCCCTGCGCC         1,00         0,37 Hs.183202 ESTs           1151 GTGAATGCACT         1,00         0,37 Hs.17401 KIAA0699 protein	1137	TCTTTAAAGTA	1,00	0,37	Hs.25155	
1139 GTGGCGAATGC         1,00         0,37 Hs.230479 EST           1140 TATTAACATTC         1,00         0,37 Hs.226573 inhibitor of kappa light polypeptide gene enhancer i           1141 TCTCCATTCCT         1,00         0,37 Hs.226573 inhibitor of kappa light polypeptide gene enhancer i           1142 TATATAGAATG         1,00         0,37 Hs.22581 ESTs           1143 GCGAGATCCTG         1,00         0,37 Hs.22529 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1144 CACTCCAACCT         1,00         0,37 Hs.205899 ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC           1145 CGATGTTAAAA         1,00         0,37 Hs.20072 myosin regulatory light chain interacting protein           1146 CCTTCTGAATA         1,00         0,37 Hs.194660 ceroid-lipofuscinosis, neuronal 3, juvenil (Batten,           1147 TGAGGATCCAG         1,00         0,37 Hs.191621 ESTs           1148 ATTGTACAACA         1,00         0,37 Hs.184326 CDC10 (cell division cycle 10, S. cerevisiae, homolo           1149 AGCCTATTAAA         1,00         0,37 Hs.183593 zinc finger protein 24 (KOX 17)           1150 GCCCCTGCGCC         1,00         0,37 Hs.183202 ESTs           1151 GTGAATGCACT         1,00         0,37 Hs.17401 KIAA0699 protein	1138	ATATTGGTGGT	1,00	0,37	Hs.250692	
enhancer i	1139	GTGGCGAATGC	1,00			
enhancer i	1140	TATTAACATTC	1,00	0,37	Hs.226573	
1142         TATATAGAATG         1,00         0,37         Hs.22581         ESTs           1143         GCGAGATCCTG         1,00         0,37         Hs.22529         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1144         CACTCCAACCT         1,00         0,37         Hs.205899         ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC           1145         CGATGTTAAAA         1,00         0,37         Hs.20072         myosin regulatory light chain interacting protein           1146         CCTTCTGAATA         1,00         0,37         Hs.194660         ceroid-lipofuscinosis, neuronal 3, juvenil (Batten,           1147         TGAGGATCCAG         1,00         0,37         Hs.184326         CDC10 (cell division cycle 10, S. cerevisiae, homolo           1149         AGCCTATTAAA         1,00         0,37         Hs.183593         zinc finger protein 24 (KOX 17)           1150         GCCCCTGCGCC         1,00         0,37         Hs.183202         ESTs           1151         GTGAATGCACT         1,00         0,37         Hs.17411         KIAA0699 protein	1141	TCTCCATTCCT	1,00	0,37	Hs.226573	
1143         GCGAGATCCTG         1,00         0,37         Hs.22529         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1144         CACTCCAACCT         1,00         0,37         Hs.205899         ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC           1145         CGATGTTAAAA         1,00         0,37         Hs.20072         myosin regulatory light chain interacting protein           1146         CCTTCTGAATA         1,00         0,37         Hs.194660         ceroid-lipofuscinosis, neuronal 3, juvenil (Batten,           1147         TGAGGATCCAG         1,00         0,37         Hs.191621         ESTs           1148         ATTGTACAACA         1,00         0,37         Hs.184326         CDC10 (cell division cycle 10, S. cerevisiae, homolo           1149         AGCCTATTAAA         1,00         0,37         Hs.183593 zinc finger protein 24 (KOX 17)           1150         GCCCCTGCGCC         1,00         0,37         Hs.183202         ESTs           1151         GTGAATGCACT         1,00         0,37         Hs.17411         KIAA0699 protein	1142	TATATAGAATG	1,00	0,37	Hs.22581	ESTs
ALU SUBFAMILY SC	1143	GCGAGATCCTG		0,37	Hs.22529	
protein						ALU SUBFAMILY SC
Catten,   1147 TGAGGATCCAG   1,00   0,37 Hs.191621 ESTs   1148 ATTGTACAACA   1,00   0,37 Hs.184326 CDC10 (cell division cycle 10, S. cerevisiae, homolo   1149 AGCCTATTAAA   1,00   0,37 Hs.183593 zinc finger protein 24 (KOX 17)   1150 GCCCTGCGCC   1,00   0,37 Hs.183202 ESTs   1151 GTGAATGCACT   1,00   0,37 Hs.17411 KIAA0699 protein   1152 TTCATTAAGAA   1,00   0,37 Hs.17411 KIAA0699 protein   1147 KIAA0699 protein   1147 KIAA0699 protein   1148 KIAA0699 protein   1149 KIAA0699 prot						protein
1148       ATTGTACAACA       1,00       0,37       Hs.184326       CDC10 (cell division cycle 10, S. cerevisiae, homolo         1149       AGCCTATTAAA       1,00       0,37       Hs.183593 zinc finger protein 24 (KOX 17)         1150       GCCCCTGCGCC       1,00       0,37       Hs.183202       ESTs         1151       GTGAATGCACT       1,00       0,37       Hs.176065       ESTs         1152       TTCATTAAGAA       1,00       0,37       Hs.17411       KIAA0699 protein	1146	CCTTCTGAATA	1,00	0,37	Hs.194660	, · · · · · · · · · · · · · · · · · · ·
cerevisiae, homolo   1149 AGCCTATTAAA   1,00   0,37 Hs.183593 zinc finger protein 24 (KOX 17)   1150 GCCCTGCGCC   1,00   0,37 Hs.183202 ESTs   1151 GTGAATGCACT   1,00   0,37 Hs.176065 ESTs   1152 TTCATTAAGAA   1,00   0,37 Hs.17411 KIAA0699 protein   1152 KIAA0699 protein   1152 KIAA0699 protein   1153 KIAA0699 protein   1154 KIAA0699 protein   1154 KIAA0699 protein   1155 KIAA0699 prot						
1150 GCCCTGCGCC       1,00       0,37 Hs.183202 ESTs         1151 GTGAATGCACT       1,00       0,37 Hs.176065 ESTs         1152 TTCATTAAGAA       1,00       0,37 Hs.17411 KIAA0699 protein	1148	ATTGTACAACA	1,00	0,37	Hs.184326	
1151 GTGAATGCACT         1,00         0,37 Hs.176065 ESTs           1152 TTCATTAAGAA         1,00         0,37 Hs.17411 KIAA0699 protein				0,37	Hs.183593	zinc finger protein 24 (KOX 17)
1152 TTCATTAAGAA 1,00 0,37 Hs.17411 KIAA0699 protein				0,37	Hs.183202	ESTs
			1,00			
1153 ACCGAGGTGCA 1,00 0,37 Hs.171882 ESTs	1152	TTCATTAAGAA	1,00	0,37	Hs.17411	KIAA0699 protein
			1,00			
1154 TTGGTATTGCA 1,00 0,37 Hs.163541 ESTs	1154	TTGGTATTGCA	1,00	0,37	Hs.163541	ESTs

1155   GTGAATTTTCA	4455	ATOTATTTOA	4 00	0.07	11- 404554	It
NT2RP2000845     NT2RP2000845   NT2RP2000845   NT2RP2000845     NT2RP2000845   NT2RP2000845     NT2RP2000845     NT2RP2000845   NT2RP2000845   NT2RP2000845   NT2RP2000845   NT2RP2000845   NT2RP2000845   NT2RP						
1158   GTGGTGCAAAC	1156	GIIGAAIIGCA	1,00			NT2RP2000845
ALU SUBFAMILY J SE			1,00			
1160   TTTGCTTTTGA	1158	GTGGTGCAAAC	1,00	0,37	Hs.149852	
1161   AAACCATTAGA	1159	GGCTTTGGAAT	1,00	0,37	Hs.146481	ESTs
1162   AAACACCCCA	1160	TTTGCTTTTGA	1,00	0,37	Hs.144504	hypothetical protein FLJ10624
CD55, Crom	1161	AAAGCATTAGA	1,00	0,37	Hs.14155	KIAA0653 protein
1164   GTGCTGCACAC	1162	AAACAACCCCA	1,00	0,37	Hs.1369	
IIII ALU CLASS C   1165 TTTAAACTTGG	1163	CTCCTTGTCCC	1,00			
1166   CGGCCCAGGTT	1164	GTGCTGCACAC	1,00	0,37		IIII ALU CLASS C
International	1165	TTTAAACTTGG	1,00			
International	1166	CGGCCCAGGTT		0,37	Hs.122823	thousand and one amino acid protein
1168 ATAGATACACA						kinase
1169   TGGCAGTAGTG				0,37	Hs.121509	collagen, type XI, alpha 2
1170	-					
1172 TACACCTGGAA         1,00         0,37 Hs.114624 ESTs           1173 CAAGGATTTT         1,00         0,37 Hs.11323 Protein inhibitor of activated STAT X           1174 CTAGTATAAAA         1,00         0,37 Hs.106650 hypothetical protein FLJ20533           1175 AATATAAAAAA         1,00         0,37 Hs.103548 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1176 CAAATATGGTT         1,00         0,37 Hs.10351 KIAA0308 protein           1177 CAAGAACAGGG         1,00         0,37 Hs.102135 signal sequence receptor, delta (translocon-associat           1178 TCACCGGTCAG         61,00         16,20 Hs.80562 gelsolin (amyloidosis, Finnish type)           1179 GTGAAAACCCC         4,00         1,38 Hs.277213 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180 GTGGTGGGCAC         60,00         15,85 Hs.77510 isovaleryl Coenzyme A dehydrogenase           1181 GTGGTGGACC         60,00         15,81 Hs.136509 EST           1183 CATCACGGATC         3,00         1,08 Hs.278329 ESTs           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibit	1170	AAATCCTTCTA	1,00			(annexin II ligand,
1172 TACACCTGGAA         1,00         0,37 Hs.114624 ESTs           1173 CAAGGATTTT         1,00         0,37 Hs.11323 Protein inhibitor of activated STAT X           1174 CTAGTATAAAA         1,00         0,37 Hs.106650 hypothetical protein FLJ20533           1175 AATATAAAAAA         1,00         0,37 Hs.103548 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1176 CAAATATGGTT         1,00         0,37 Hs.10351 KIAA0308 protein           1177 CAAGAACAGGG         1,00         0,37 Hs.102135 signal sequence receptor, delta (translocon-associat           1178 TCACCGGTCAG         61,00         16,20 Hs.80562 gelsolin (amyloidosis, Finnish type)           1179 GTGAAAACCCC         4,00         1,38 Hs.277213 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180 GTGGTGGGCAC         60,00         15,85 Hs.77510 isovaleryl Coenzyme A dehydrogenase           1181 GTGGTGGACC         60,00         15,81 Hs.136509 EST           1183 CATCACGGATC         3,00         1,08 Hs.278329 ESTs           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibit	1171	TCAAACTTTGT	1,00	0,37	Hs.117582	CGI-43 protein
1173 CAAGGATTTTT         1,00         0,37 Hs.111323 Protein inhibitor of activated STAT X           1174 CTAGTATAAAA         1,00         0,37 Hs.106650 hypothetical protein FLJ20533           1175 AATATAAAAAA         1,00         0,37 Hs.103548 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1176 CAAATATGGTT         1,00         0,37 Hs.10351 KIAA0308 protein           1177 CAAGAACAGGG         1,00         0,37 Hs.102135 signal sequence receptor, delta (translocon-associat (translocon-associat (translocon-associat))           1178 TCACCGGTCAG         61,00         16,20 Hs.80562 gelsolin (amyloidosis, Finnish type)           1179 GTGAAAACCCC         4,00         1,38 Hs.277213 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180 GTGGTGGGCAC         60,00         15,85 Hs.77510 isovaleryl Coenzyme A dehydrogenase           1181 GTGGTGGGTGC         60,00         15,81 Hs.136509 EST           1182 CCGTTGCACTC         6,00         1,96 Hs.278329 ESTs           1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.13766 ESTs           1185 CCCTTTATATC         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00	1172	TACACCTGGAA	1,00	0,37	Hs.114624	ESTs
1174 CTAGTATAAAA         1,00         0,37 Hs.106650 hypothetical protein FLJ20533           1175 AATATAAAAAA         1,00         0,37 Hs.103548 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1176 CAAATATGGTT         1,00         0,37 Hs.10351 KIAA0308 protein           1177 CAAGAACAGGG         1,00         0,37 Hs.102135 signal sequence receptor, delta (translocon-associat (translocon-associat)           1178 TCACCGGTCAG         61,00         16,20 Hs.80562 gelsolin (amyloidosis, Finnish type)           1179 GTGAAAACCCC         4,00         1,38 Hs.277213 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180 GTGGTGGGCAC         60,00         15,85 Hs.77510 isovaleryl Coenzyme A dehydrogenase           1181 GTGGTGGGTGC         60,00         15,81 Hs.136509 EST           1182 CCGTTGCACTC         6,00         1,96 Hs.278329 ESTs           1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG	1173	CAAGGATTTTT	1,00			
1175         AATATAAAAAA         1,00         0,37         Hs.103548         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           1176         CAAATATGGTT         1,00         0,37         Hs.10351         KIAA0308 protein           1177         CAAGAACAGGG         1,00         0,37         Hs.102135         signal sequence receptor, delta (translocon-associat (translocon-associat)           1178         TCACCGGTCAG         61,00         16,20         Hs.80562         gelsolin (amyloidosis, Finnish type)           1179         GTGAAAACCCC         4,00         1,38         Hs.277213         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180         GTGGTGGGCAC         60,00         15,85         Hs.77510         isovaleryl Coenzyme A dehydrogenase           1181         GTGGTGGGTCC         60,00         15,81         Hs.136509         EST           1182         CCGTTGCACTC         6,00         1,96         Hs.278329         ESTs           1183         CATCACGGATC         3,00         1,08         Hs.82112         interleukin 1 receptor, type I           1184         GTATGTACAGG         3,00         1,08         Hs.13766         ESTs           1185         CCCTTTATATC         3,00         1,08         Hs.106070         cycl	1174	CTAGTATAAAA				
1176 CAAATATGGTT         1,00         0,37 Hs.10351         KIAA0308 protein           1177 CAAGAACAGGG         1,00         0,37 Hs.102135         signal sequence receptor, delta (translocon-associat           1178 TCACCGGTCAG         61,00         16,20 Hs.80562         gelsolin (amyloidosis, Finnish type)           1179 GTGAAAACCCC         4,00         1,38 Hs.277213         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180 GTGGTGGGCAC         60,00         15,85 Hs.77510         isovaleryl Coenzyme A dehydrogenase           1181 GTGGTGGGTGC         60,00         15,81 Hs.136509         EST           1182 CCGTTGCACTC         6,00         1,96 Hs.278329         ESTs           1183 CATCACGGATC         3,00         1,08 Hs.82112         interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255         ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.119475         cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070         cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841         ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG         5,00         1,68 Hs.920         modulator recognition	1175	AATATAAAAAA				ESTs, Weakly similar to ALU1_HUMAN
1177         CAAGAACAGGG         1,00         0,37         Hs.102135         signal sequence receptor, delta (translocon-associat           1178         TCACCGGTCAG         61,00         16,20         Hs.80562         gelsolin (amyloidosis, Finnish type)           1179         GTGAAAACCCC         4,00         1,38         Hs.277213         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180         GTGGTGGGCAC         60,00         15,85         Hs.77510         isovaleryl Coenzyme A dehydrogenase           1181         GTGGTGGGTGC         60,00         15,81         Hs.136509         EST           1182         CCGTTGCACTC         6,00         1,96         Hs.278329         ESTs           1183         CATCACGGATC         3,00         1,08         Hs.82112         interleukin 1 receptor, type I           1184         GTATGTACAGG         3,00         1,08         Hs.164255         ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185         CCCTTTATATC         3,00         1,08         Hs.19475         cold inducible RNA-binding protein           1187         CCCATCTAGCT         3,00         1,08         Hs.106070         cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188         TCTCAAAAAAAA         8,00         2,52	1176	CAAATATGGTT	1,00	0,37	Hs.10351	KIAA0308 protein
1178 TCACCGGTCAG         61,00         16,20 Hs.80562         gelsolin (amyloidosis, Finnish type)           1179 GTGAAAACCCC         4,00         1,38 Hs.277213 EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180 GTGGTGGGCAC         60,00         15,85 Hs.77510 isovaleryl Coenzyme A dehydrogenase           1181 GTGGTGGGTGC         60,00         15,81 Hs.136509 EST           1182 CCGTTGCACTC         6,00         1,96 Hs.278329 ESTs           1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG         5,00         1,68 Hs.920 modulator recognition factor I           1190 GGTTATTTAGT         5,00         1,68 Hs.8110 adducin 3 (gamma)	1177	CAAGAACAGGG		0,37	Hs.102135	signal sequence receptor, delta
1179         GTGAAAACCCC         4,00         1,38         Hs.277213         EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ S           1180         GTGGTGGGCAC         60,00         15,85         Hs.77510         isovaleryl Coenzyme A dehydrogenase           1181         GTGGTGGGTGC         60,00         15,81         Hs.136509         EST           1182         CCGTTGCACTC         6,00         1,96         Hs.278329         ESTs           1183         CATCACGGATC         3,00         1,08         Hs.82112         interleukin 1 receptor, type I           1184         GTATGTACAGG         3,00         1,08         Hs.164255         ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185         CCCTTTATATC         3,00         1,08         Hs.13766         ESTs           1186         GCTCGTGGTCA         3,00         1,08         Hs.119475         cold inducible RNA-binding protein           1187         CCCATCTAGCT         3,00         1,08         Hs.106070         cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188         TCTCAAAAAAA         8,00         2,52         Hs.194841         ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189         CCCCTGGCTGG         5,00         1,68         Hs.920 <td< td=""><td>1178</td><td>TCACCGGTCAG</td><td>61,00</td><td>16,20</td><td>Hs.80562</td><td>gelsolin (amyloidosis, Finnish type)</td></td<>	1178	TCACCGGTCAG	61,00	16,20	Hs.80562	gelsolin (amyloidosis, Finnish type)
1181 GTGGTGGGTGC         60,00         15,81 Hs.136509 EST           1182 CCGTTGCACTC         6,00         1,96 Hs.278329 ESTs           1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2 HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841 ESTs, Moderately similar to ALU8 HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG         5,00         1,68 Hs.920 modulator recognition factor I           1190 GGTTATTTAGT         5,00         1,68 Hs.8110 adducin 3 (gamma)	1179	GTGAAAACCCC	4,00	1,38	Hs.277213	EST, Weakly similar to ALU7_HUMAN
1181 GTGGTGGGTGC         60,00         15,81 Hs.136509 EST           1182 CCGTTGCACTC         6,00         1,96 Hs.278329 ESTs           1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2 HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841 ESTs, Moderately similar to ALU8 HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG         5,00         1,68 Hs.920 modulator recognition factor I           1190 GGTTATTTAGT         5,00         1,68 Hs.8110 adducin 3 (gamma)				15,85	Hs.77510	isovaleryl Coenzyme A dehydrogenase
1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG         5,00         1,68 Hs.920 modulator recognition factor I           1190 GGTTATTTAGT         5,00         1,68 Hs.8110 adducin 3 (gamma)				15,81	Hs.136509	EST
1183 CATCACGGATC         3,00         1,08 Hs.82112 interleukin 1 receptor, type I           1184 GTATGTACAGG         3,00         1,08 Hs.164255 ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185 CCCTTTATATC         3,00         1,08 Hs.13766 ESTs           1186 GCTCGTGGTCA         3,00         1,08 Hs.119475 cold inducible RNA-binding protein           1187 CCCATCTAGCT         3,00         1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188 TCTCAAAAAAA         8,00         2,52 Hs.194841 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189 CCCCTGGCTGG         5,00         1,68 Hs.920 modulator recognition factor I           1190 GGTTATTTAGT         5,00         1,68 Hs.8110 adducin 3 (gamma)	1182	CCGTTGCACTC	6,00	1,96	Hs.278329	ESTs
1184         GTATGTACAGG         3,00         1,08         Hs. 164255         ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY           1185         CCCTTTATATC         3,00         1,08         Hs. 13766         ESTs           1186         GCTCGTGGTCA         3,00         1,08         Hs. 119475         cold inducible RNA-binding protein           1187         CCCATCTAGCT         3,00         1,08         Hs. 106070         cyclin-dependent kinase inhibitor 1C (p57, Kip2)           1188         TCTCAAAAAAA         8,00         2,52         Hs. 194841         ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY           1189         CCCCTGGCTGG         5,00         1,68         Hs. 920         modulator recognition factor I           1190         GGTTATTTAGT         5,00         1,68         Hs. 8110         adducin 3 (gamma)			3,00			
1186 GCTCGTGGTCA3,001,08 Hs.119475 cold inducible RNA-binding protein1187 CCCATCTAGCT3,001,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)1188 TCTCAAAAAAA8,002,52 Hs.194841 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY1189 CCCCTGGCTGG5,001,68 Hs.920 modulator recognition factor I1190 GGTTATTTAGT5,001,68 Hs.8110 adducin 3 (gamma)	1184	GTATGTACAGG	3,00	1,08		
1187 CCCATCTAGCT 3,00 1,08 Hs.106070 cyclin-dependent kinase inhibitor 1C (p57, Kip2)  1188 TCTCAAAAAAA 8,00 2,52 Hs.194841 ESTs, Moderately similar to ALU8 HUMAN ALU SUBFAMILY  1189 CCCCTGGCTGG 5,00 1,68 Hs.920 modulator recognition factor I  1190 GGTTATTTAGT 5,00 1,68 Hs.8110 adducin 3 (gamma)	1185	CCCTTTATATC	3,00	1,08		
(p57, Kip2)   1188   TCTCAAAAAAA	1186	GCTCGTGGTCA	3,00	1,08	Hs.119475	cold inducible RNA-binding protein
1188 TCTCAAAAAAA 8,00 2,52 Hs.194841 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY  1189 CCCCTGGCTGG 5,00 1,68 Hs.920 modulator recognition factor I  1190 GGTTATTTAGT 5,00 1,68 Hs.8110 adducin 3 (gamma)			3,00	1,08		•
1189 CCCCTGGCTGG5,001,68 Hs.920modulator recognition factor I1190 GGTTATTTAGT5,001,68 Hs.8110adducin 3 (gamma)	1188	TCTCAAAAAA	8,00	2,52		ESTs, Moderately similar to
1190 GGTTATTTAGT 5,00 1,68 Hs.8110 adducin 3 (gamma)	1189	CCCCTGGCTGG	5,00	1,68	Hs.920	
i i			7,00			

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1122					ALU SUBFAMILY SQ
	AGTTGTTTGGT	2,00			ESTs
	TCATAGCCTTG	2,00			heat shock 27kD protein 2
	AGGACTGGACT	2,00			H2A histone family, member Y
	TAAACCTAGGA	2,00			EGF-like-domain, multiple 3
$\overline{}$	GTGGCTCACTT	2,00		Hs.285616	
	TCATTTGGTGT	2,00		Hs.285439	
	GCCTTGGCAGT	2,00			iroquois-class homeodomain protein
1199	CCCTTGTTCTT	2,00	0,75	Hs.250723	FK506 binding protein 12-rapamycin
L					associated protei
	GAACAGTATGA	2,00		Hs.189762	
1201	ATGGCAGGCGG	2,00	0,75	Hs.161554	hypothetical protein FLJ20159
1202	ACACAGCAAGA	136,00	34,48	Hs.80562	gelsolin (amyloidosis, Finnish type)
1203	GATCAGGCCAG	30,00	8,16	Hs.119571	collagen, type III, alpha 1 (Ehlers-Danios
					syndrome
1204	GTGAAACTCTG	32,00	8,57	Hs.188853	Homo sapiens cDNA FLJ10150 fis, clone
,					HEMBA1003395
1205	GGCCTGCAGGA	6,00	1,93	Hs.71869	apoptosis-associated speck-like protein
			·		containing a
1206	AAATCAATACA	7,00	2,21	Hs.94953	ESTs, Highly similar to C1QC_HUMAN
			·		COMPLEMENT C1Q SU
1207	ATTGTACCACT	7,00	2,21	Hs.7099	hypothetical protein FLJ20265
	CGCCTGTAGTC	29,00			ESTs
	GCAAAACCCAG	4,00			ESTs, Moderately similar to
1		.,,,,,	.,00		ALU6_HUMAN ALU SUBFAMILY
1210	CTTTGATGCGG	4,00	1.35	Hs.183601	regulator of G-protein signalling 16
	GGCCCTAGGCA	33,00		Hs.78909	butyrate response factor 2 (EGF-
		00,00	٠,, ٠		response factor 2)
1212	CCTGGCTAATT	29,00	7.72	Hs.25661	ESTs, Weakly similar to ALUF_HUMAN
			• ,• =		IIII ALU CLASS F
1213	CTTCCTGGCCT	10,00	2.96	Hs.83623	nuclear receptor subfamily 1, group I,
1	,	.0,00	_,00		member 3
1214	GCGGGGTGGAG	37,00	9.65	Hs.85155	butyrate response factor 1 (EGF-
		0,,00	5,55	110.00100	response factor 1)
1215	GTGGCAGGCGC	85,00	21.18	Hs.48604	DKFZP434B168 protein
	AGCCCAGGAGG	9,00			ESTs, Weakly similar to unnamed
1.2.0		0,00	_,00		protein product [H.s
1217	ATAGTGCCACT	6,00	1 90		ESTs, Weakly similar to ALU7_HUMAN
1		0,00	1,00	110.2-107 17	ALU SUBFAMILY SQ
1218	CATTTGTAAAA	3,00	1.05	Hs.84429	KIAA0971 protein
	CGTACAGCCCC	3,00			KIAA1448 protein
-	GGGCTACGTCC	3,00			kallikrein 1, renal/pancreas/salivary
	ATCACACCACT	27,00			KIAA0707 protein
	CACTCCAGCCT	13,00			ESTs, Weakly similar to ALU7_HUMAN
1222	OACTOCAGCCT	13,00	3,09	15. 13343	ALU SUBFAMILY SQ
1222	CTTGTAATCCC	46,00	11 60	Uc 192252	ESTs, Weakly similar to ALU1 HUMAN
1223	CHGIMAICCC	40,00	11,09	⊓S. 103233 	ALU SUBFAMILY J S
1224	CTCAAACCTCA	47.00	11.04	Un 152020	<u> </u>
$\overline{}$	GTGAAACCTCA	47,00		Hs.153029	
1225	ATCTCAGCTCA	12,00	3,43	<sub>[HS.246192</sub>	ESTs, Weakly similar to RMS1_HUMAN

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1227   ACTCGAATATC	
1228   GAGTCCCTGGT	
1229   TGCAATATGCC   6,00   1,87   Hs.750   fibrillin 1 (Marfan syndrome)     1230   GAAGCAATAAA   6,00   1,87   Hs.198253   major histocompatibility complex, c     1, DQ alpha   ESTs, Moderately similar to     1, DQ alpha   ESTs, Moderately similar	
1230   GAAGCAATAAA   6,00   1,87   Hs.198253   major histocompatibility complex, c   II, DQ alpha   1231   GTGAAACTCCG   37,00   9,36   Hs.261734   ESTs, Moderately similar to ALUT_HUMAN ALU SUBFAMILY   1232   GGCTGCCGAGT   2,00   0,72   Hs.99829   hypothetical protein FLJ20565   1233   CTTAGCCCCAG   2,00   0,72   Hs.96908   ESTs   1234   TTATTCCACAA   2,00   0,72   Hs.9619   SRY (sex determining region Y)-bo   1236   TTTCCTTTTG   2,00   0,72   Hs.78546   ATPase, Ca++ transporting, plasma membrane 1   1237   CTTGCATAAGA   2,00   0,72   Hs.78546   ATPase, Ca++ transporting, plasma membrane 1   1238   TATGTGTTCT   2,00   0,72   Hs.3353   beta-1,3-glucuronyltransferase 1   (glucuronosyltransf   1239   ACTATCATCTT   2,00   0,72   Hs.29117   H.sapiens mRNA for pur alpha extermining region Y)-bo   1239   ACTATCATCTT   2,00   0,72   Hs.271166   ESTs, Moderately similar to   1240   CGCCTATAGTC   2,00   0,72   Hs.271166   ESTs, Moderately similar to   1241   CCCGCCAGTGC   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy   1242   CGGCAGTGC   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy   1243   CGGGAAGACAT   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy   1244   CCAGTAGTCCC   2,00   0,72   Hs.147959   EST   1245   CAGTITGTTT   2,00   0,72   Hs.144077   hypothetical protein PRO2975   1246   TCCTTAAAAT   2,00   0,72   Hs.144477   hypothetical protein PRO2975   1248   AGAATCACTTG   41,00   1,61   Hs.117582   CGI-43 protein   1248   AGAATCACTTG   41,00   1,016   Hs.117582   CGI-43 protein   1250   GCCCCAGAATC   3,00   1,02   Hs.18269   ESTs, Moderately similar to   1,29   Hs.232598   CREB binding protein (Rubinstein-1)   1,29   Hs.232598   CREB binding protein (Rubinstein-1)   1,29   Hs.232540   ESTs   1,29   Hs.232540   E	
II, ĎQ alpha	
ALUT_HUMAN ALU SUBFAMILY	lass
1233   CTTAGCCCAG	
1234   TTATTCCACAA   2,00   0,72   Hs.93765   lipoma HMGIC fusion partner   1235   TCACAGCCCCC   2,00   0,72   Hs.8619   SRY (sex determining region Y)-bo   1236   TTTCCTTTTG   2,00   0,72   Hs.78546   ATPase, Ca++ transporting, plasma membrane   1237   CTTGCATAAGA   2,00   0,72   Hs.72912   cytochrome P450, subfamily   (aron compound-indu   1238   TATGTGTTCTC   2,00   0,72   Hs.3353   beta-1,3-glucuronyltransferase 1   (glucuronosyltransf   1239   ACTATCATCTT   2,00   0,72   Hs.29117   H.sapiens mRNA for pur alpha exterminated   1240   CGCCTATAGTC   2,00   0,72   Hs.271166   ESTs, Moderately similar to ALUT   HUMAN ALU SUBFAMILY   1241   CCCGCCAGTGC   2,00   0,72   Hs.256297   integrin, alpha 11   1242   TTCTAATTTTT   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy   1243   CGGGAAGACAT   2,00   0,72   Hs.154525   KIAA1076 protein   1244   CCAGTAGTCC   2,00   0,72   Hs.14477   hypothetical protein PRO2975   1246   TCCTTTAAAAT   2,00   0,72   Hs.14477   hypothetical protein PRO2975   1246   TCCTTTAAAAT   2,00   0,72   Hs.10587   KIAA0353 protein   1247   GTGCTAAGCGG   46,00   11,41   Hs.4217   collagen, type VI, alpha 2   1248   AGAATCACTTG   41,00   10,16   Hs.117582   CGI-43 protein   1249   GTGGTAAGCG   40,00   1,02   Hs.8682   ESTs, Moderately similar to ALU1   HUMAN ALU SUBFAMILY   1251   TTGGGAGGCTG   3,00   1,02   Hs.18269   ESTs, Weakly similar to A46010 X-retinopathy   1252   GTGGCACGCGC   19,00   5,01   Hs.187346   ESTs   1,29   Hs.23598   CREB binding protein (Rubinstein-1 syndrome)   1254   AGTTCGAGACC   4,00   1,29   Hs.23594   ESTs   1254   AGTTCGAGACC   4,00   1,29   Hs.235540   ESTs   1254   AGTTCGAGACC   4,00   1,29   Hs.235540   ESTs   1254   AGTTCGAGACC   4,00   1,29   Hs.235540   ESTs   1255   CAAGCGCTCTA   4,00   1,29   Hs.235540   ESTs   1254   AGTTCGAGACC   4,00   1,29   Hs.235540   ESTs   1255   CAAGCGCTCTA   4,00	
1235   TCACAGCCCCC   2,00   0,72   Hs.8619   SRY (sex determining region Y)-bo   1236   TTTCCTTTTG   2,00   0,72   Hs.78546   ATPase, Ca++ transporting, plasma membrane 1   237   CTTGCATAAGA   2,00   0,72   Hs.72912   cytochrome P450, subfamily I (aron compound-indu   1238   TATGTGTTCTC   2,00   0,72   Hs.3353   beta-1,3-glucuronyltransferase 1 (glucuronosyltransf   1239   ACTATCATCTT   2,00   0,72   Hs.29117   H.sapiens mRNA for pur alpha extermal substantial of the state of the	
1235   TCACAGCCCCC   2,00   0,72   Hs.8619   SRY (sex determining region Y)-bo   1236   TTTCCTTTTG   2,00   0,72   Hs.78546   ATPase, Ca++ transporting, plasma membrane 1   237   CTTGCATAAGA   2,00   0,72   Hs.72912   cytochrome P450, subfamily I (aron compound-indu   1238   TATGTGTTCTC   2,00   0,72   Hs.3353   beta-1,3-glucuronyltransferase 1 (glucuronosyltransf   1239   ACTATCATCTT   2,00   0,72   Hs.29117   H.sapiens mRNA for pur alpha extermal substantial of the state of the	
1236   TTTCCTTTTG	x 18
Compound-indu     1238 TATGTGTTCTC   2,00   0,72   Hs.3353   beta-1,3-glucuronyltransferase 1     1239 ACTATCATCTT   2,00   0,72   Hs.29117   H.sapiens mRNA for pur alpha exteraction     1240 CGCCTATAGTC   2,00   0,72   Hs.271166   ESTs, Moderately similar to     1241 CCCGCCAGTGC   2,00   0,72   Hs.256297   integrin, alpha 11     1242 TTCTAATTTT   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy     1243 CGGGAAGACAT   2,00   0,72   Hs.154525   KIAA1076 protein     1244 CCAGTAGTCCC   2,00   0,72   Hs.144959   EST     1245 CAGTTTGTGTT   2,00   0,72   Hs.144477   hypothetical protein PRO2975     1246 TCCTTTAAAAT   2,00   0,72   Hs.10587   KIAA0353 protein     1247 GTGCTAAGCGG   46,00   11,41   Hs.4217   collagen, type VI, alpha 2     1248 AGAATCACTTG   41,00   10,16   Hs.117582   CGI-43 protein     1249 GTGGTGTACGC   9,00   2,60   Hs.182225   RNA binding motif protein 3     1250 GCCCCAGAATC   3,00   1,02   Hs.8682   ESTs, Moderately similar to     ALU1 HUMAN ALU SUBFAMILY     1251 TTGGGAGGCTG   3,00   1,02   Hs.118269   ESTs, Weakly similar to     1252 GTGGCACGCGC   19,00   5,01   Hs.187346   ESTs     1253 CAAGCGCTCTA   4,00   1,29   Hs.23598   CREB binding protein (Rubinstein-1 syndrome)     1254 AGTTCGAGACC   4,00   1,29   Hs.232540   ESTs	
(glucuronosyltransf   1239   ACTATCATCTT   2,00   0,72   Hs.29117   H.sapiens mRNA for pur alpha exter 3'untranslated   1240   CGCCTATAGTC   2,00   0,72   Hs.271166   ESTs, Moderately similar to ALUT_HUMAN ALU SUBFAMILY   1241   CCCGCCAGTGC   2,00   0,72   Hs.256297   integrin, alpha 11   1242   TTCTAATTTTT   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy 4   1243   CGGGAAGACAT   2,00   0,72   Hs.154525   KIAA1076   protein   1244   CCAGTAGTCCC   2,00   0,72   Hs.147959   EST   1245   CAGTTGTGTT   2,00   0,72   Hs.14477   hypothetical protein PRO2975   1246   TCCTTTAAAAT   2,00   0,72   Hs.10587   KIAA0353   protein   1247   GTGCTAAGCGG   46,00   11,41   Hs.4217   collagen, type VI, alpha 2   1248   AGAATCACTTG   41,00   10,16   Hs.117582   CGI-43   protein   1249   GTGGTGTACGC   9,00   2,60   Hs.182225   RNA binding motif protein 3   1250   GCCCCAGAATC   3,00   1,02   Hs.8682   ESTs, Moderately similar to ALU1   HUMAN ALU SUBFAMILY   1251   TTGGGAGGCTG   3,00   1,02   Hs.118269   ESTs, Weakly similar to A46010 X-retinopathy   1252   GTGGCACGCGC   19,00   5,01   Hs.187346   ESTs   1253   CAAGCGCTCTA   4,00   1,29   Hs.23598   CREB binding protein (Rubinstein-Tsyndrome)   1254   AGTTCGAGACC   4,00   1,29   Hs.232540   ESTs   1255	natic
3'untranslated   1240   CGCCTATAGTC   2,00   0,72   Hs.271166   ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY   1241   CCCGCCAGTGC   2,00   0,72   Hs.256297   integrin, alpha 11   1242   TTCTAATTTTT   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy 4   1243   CGGGAAGACAT   2,00   0,72   Hs.154525   KIAA1076 protein   1244   CCAGTAGTCCC   2,00   0,72   Hs.147959   EST   1245   CAGTTTGTGTT   2,00   0,72   Hs.14477   hypothetical protein PRO2975   1246   TCCTTTAAAAT   2,00   0,72   Hs.10587   KIAA0353 protein   1247   GTGCTAAGCGG   46,00   11,41   Hs.4217   collagen, type VI, alpha 2   1248   AGAATCACTTG   41,00   10,16   Hs.117582   CGI-43 protein   1249   GTGGTGTACGC   9,00   2,60   Hs.182225   RNA binding motif protein 3   1250   GCCCCAGAATC   3,00   1,02   Hs.8682   ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY   1251   TTGGGAGGCTG   3,00   1,02   Hs.118269   ESTs, Weakly similar to A46010 X-retinopathy   1252   GTGGCACGCGC   19,00   5,01   Hs.187346   ESTs   1253   CAAGCGCTCTA   4,00   1,29   Hs.23598   CREB binding protein (Rubinstein-Tsyndrome)   1254   AGTTCGAGACC   4,00   1,29   Hs.232540   ESTs   1255   1256   1257	
ALU7_HUMAN ALU SUBFAMILY   1241   CCCGCCAGTGC   2,00   0,72   Hs.256297   integrin, alpha 11   1242   TTCTAATTTTT   2,00   0,72   Hs.170414   paired basic amino acid cleaving sy 4   1243   CGGGAAGACAT   2,00   0,72   Hs.154525   KIAA1076   protein   1244   CCAGTAGTCCC   2,00   0,72   Hs.147959   EST   1245   CAGTTTGTGTT   2,00   0,72   Hs.144477   hypothetical protein PRO2975   1246   TCCTTTAAAAT   2,00   0,72   Hs.10587   KIAA0353   protein   1247   GTGCTAAGCGG   46,00   11,41   Hs.4217   collagen, type VI, alpha 2   1248   AGAATCACTTG   41,00   10,16   Hs.117582   CGI-43   protein   1249   GTGGTGTACGC   9,00   2,60   Hs.182225   RNA binding motif protein 3   1250   GCCCCAGAATC   3,00   1,02   Hs.8682   ESTs, Moderately similar to ALU1   HUMAN ALU SUBFAMILY   1251   TTGGGAGGCTG   3,00   1,02   Hs.118269   ESTs, Weakly similar to A46010 X-retinopathy   1252   GTGGCACGCGC   19,00   5,01   Hs.187346   ESTs   1253   CAAGCGCTCTA   4,00   1,29   Hs.23598   CREB binding protein (Rubinstein-1 syndrome)   1254   AGTTCGAGACC   4,00   1,29   Hs.232540   ESTs   1255   1256   1257   125	nded
1242         TTCTAATTTTT         2,00         0,72         Hs.170414         paired basic amino acid cleaving sy 4           1243         CGGGAAGACAT         2,00         0,72         Hs.154525         KIAA1076 protein           1244         CCAGTAGTCCC         2,00         0,72         Hs.147959         EST           1245         CAGTITGTGTT         2,00         0,72         Hs.144477         hypothetical protein PRO2975           1246         TCCTITAAAAT         2,00         0,72         Hs.10587         KIAA0353 protein           1247         GTGCTAAGCGG         46,00         11,41         Hs.4217         collagen, type VI, alpha 2           1248         AGAATCACTTG         41,00         10,16         Hs.117582         CGI-43 protein           1249         GTGGTGTACGC         9,00         2,60         Hs.182225         RNA binding motif protein 3           1250         GCCCCAGAATC         3,00         1,02         Hs.8682         ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           1251         TTGGGAGGCTG         3,00         1,02         Hs.118269         ESTs           1252         GTGGCACGCGC         19,00         5,01         Hs.187346         ESTs           1253         CAAGCGCTCTA	
1243 CGGGAAGACAT   2,00   0,72   Hs.154525   KIAA1076 protein     1244 CCAGTAGTCCC   2,00   0,72   Hs.147959   EST     1245 CAGTTTGTGTT   2,00   0,72   Hs.144477   hypothetical protein PRO2975     1246 TCCTTTAAAAT   2,00   0,72   Hs.10587   KIAA0353 protein     1247 GTGCTAAGCGG   46,00   11,41   Hs.4217   collagen, type VI, alpha 2     1248 AGAATCACTTG   41,00   10,16   Hs.117582   CGI-43 protein     1249 GTGGTGTACGC   9,00   2,60   Hs.182225   RNA binding motif protein 3     1250 GCCCCAGAATC   3,00   1,02   Hs.8682   ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY     1251 TTGGGAGGCTG   3,00   1,02   Hs.118269   ESTs, Weakly similar to A46010 X-retinopathy     1252 GTGGCACGCGC   19,00   5,01   Hs.187346   ESTs     1253 CAAGCGCTCTA   4,00   1,29   Hs.23598   CREB binding protein (Rubinstein-Tsyndrome)     1254 AGTTCGAGACC   4,00   1,29   Hs.232540   ESTs	
1244 CCAGTAGTCCC         2,00         0,72 Hs.147959 EST           1245 CAGTTTGTGTT         2,00         0,72 Hs.144477 hypothetical protein PRO2975           1246 TCCTTTAAAAT         2,00         0,72 Hs.10587 KIAA0353 protein           1247 GTGCTAAGCGG         46,00         11,41 Hs.4217 collagen, type VI, alpha 2           1248 AGAATCACTTG         41,00         10,16 Hs.117582 CGI-43 protein           1249 GTGGTGTACGC         9,00         2,60 Hs.182225 RNA binding motif protein 3           1250 GCCCCAGAATC         3,00         1,02 Hs.8682 ESTs, Moderately similar to ALU1 HUMAN ALU SUBFAMILY           1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-T syndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	stem
1244 CCAGTAGTCCC         2,00         0,72 Hs.147959 EST           1245 CAGTTTGTGTT         2,00         0,72 Hs.144477 hypothetical protein PRO2975           1246 TCCTTTAAAAT         2,00         0,72 Hs.10587 KIAA0353 protein           1247 GTGCTAAGCGG         46,00         11,41 Hs.4217 collagen, type VI, alpha 2           1248 AGAATCACTTG         41,00         10,16 Hs.117582 CGI-43 protein           1249 GTGGTGTACGC         9,00         2,60 Hs.182225 RNA binding motif protein 3           1250 GCCCCAGAATC         3,00         1,02 Hs.8682 ESTs, Moderately similar to ALU1 HUMAN ALU SUBFAMILY           1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-1 syndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	
1245 CAGTTTGTGTT         2,00         0,72 Hs.144477 hypothetical protein PRO2975           1246 TCCTTTAAAAT         2,00         0,72 Hs.10587 KIAA0353 protein           1247 GTGCTAAGCGG         46,00         11,41 Hs.4217 collagen, type VI, alpha 2           1248 AGAATCACTTG         41,00         10,16 Hs.117582 CGI-43 protein           1249 GTGGTGTACGC         9,00         2,60 Hs.182225 RNA binding motif protein 3           1250 GCCCCAGAATC         3,00         1,02 Hs.8682 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-1 syndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	
1246 TCCTTTAAAAT         2,00         0,72 Hs.10587         KIAA0353 protein           1247 GTGCTAAGCGG         46,00         11,41 Hs.4217         collagen, type VI, alpha 2           1248 AGAATCACTTG         41,00         10,16 Hs.117582 CGI-43 protein           1249 GTGGTGTACGC         9,00         2,60 Hs.182225 RNA binding motif protein 3           1250 GCCCCAGAATC         3,00         1,02 Hs.8682 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-Tsyndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	
1248 AGAATCACTTG         41,00         10,16 Hs.117582 CGI-43 protein           1249 GTGGTGTACGC         9,00         2,60 Hs.182225 RNA binding motif protein 3           1250 GCCCCAGAATC         3,00         1,02 Hs.8682 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-T syndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	
1248 AGAATCACTTG         41,00         10,16 Hs.117582 CGI-43 protein           1249 GTGGTGTACGC         9,00         2,60 Hs.182225 RNA binding motif protein 3           1250 GCCCCAGAATC         3,00         1,02 Hs.8682 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY           1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-T syndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	
1250 GCCCAGAATC  3,00  1,02 Hs.8682  ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY  1251 TTGGGAGGCTG  3,00  1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy  1252 GTGGCACGCGC  19,00  5,01 Hs.187346 ESTs  1253 CAAGCGCTCTA  4,00  1,29 Hs.23598  CREB binding protein (Rubinstein-T syndrome)  1254 AGTTCGAGACC  4,00  1,29 Hs.232540 ESTs	
1250 GCCCAGAATC 3,00 1,02 Hs.8682 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY  1251 TTGGGAGGCTG 3,00 1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy  1252 GTGGCACGCGC 19,00 5,01 Hs.187346 ESTs  1253 CAAGCGCTCTA 4,00 1,29 Hs.23598 CREB binding protein (Rubinstein-T syndrome)  1254 AGTTCGAGACC 4,00 1,29 Hs.232540 ESTs	
1251 TTGGGAGGCTG         3,00         1,02 Hs.118269 ESTs, Weakly similar to A46010 X-retinopathy           1252 GTGGCACGCGC         19,00         5,01 Hs.187346 ESTs           1253 CAAGCGCTCTA         4,00         1,29 Hs.23598 CREB binding protein (Rubinstein-Tsyndrome)           1254 AGTTCGAGACC         4,00         1,29 Hs.232540 ESTs	
1253 CAAGCGCTCTA 4,00 1,29 Hs.23598 CREB binding protein (Rubinstein-Tsyndrome) 1254 AGTTCGAGACC 4,00 1,29 Hs.232540 ESTs	linked
1253 CAAGCGCTCTA 4,00 1,29 Hs.23598 CREB binding protein (Rubinstein-1 syndrome) 1254 AGTTCGAGACC 4,00 1,29 Hs.232540 ESTs	
1254 AGTTCGAGACC 4,00 1,29 Hs.232540 ESTs	aybi
1255 AGAACCAAAAA 4,00 1,29 Hs.181244 major histocompatibility complex, cl	ass I,
1256 CTGGCTATCCG 4,00 1,29 Hs.10784 hypothetical protein FLJ20037	
1257 TAGTCCCAGCT 11,00 3,08 Hs.274579 ancient conserved domain protein 1	
1258 GTGAAATCCTG 28,00 7,06 Hs.53531 lipoic acid synthetase	
1259 CCTGTAATTCC 47,00 11,36 Hs.23582 tumor-associated calcium signal transducer 2	
1260 CTTCTTGCCCC 20,00 5,17 Hs.251577 hemoglobin, alpha 1	

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1261	TGGTTGGTGGT	11,00	3.05	Hs.12701	plasmolipin
	CCCGTAATCCC	17,00	4.46	Hs.274168	Homo sapiens mRNA; cDNA
1.202		,55	.,		DKFZp761P0212 (from clone DK
1263	CCTGGCCAGAA	5,00	1.55	Hs.261734	ESTs, Moderately similar to
		, ,,,,	.,		ALU7 HUMAN ALU SUBFAMILY
1264	CGTGTAATCCC	12,00	3,28	Hs.187761	
1265	GCGAAACCTCA	6,00			thioredoxin-like
1266	TGGTTACAAAA	6,00	1,81	Hs.3850	Homo sapiens clone 23596 mRNA
		·	·		sequence
1267	CCACAGCACTC	7,00		Hs.273828	
1268	GTGGCACGTGC	103,00	23,64	Hs.278588	ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB
1269	ATGTCTTTTCT	18,00	4,66	Hs.1516	insulin-like growth factor-binding protein 4
1270	CTTGTAGTCCC	11,00	3,02	Hs.272202	hypothetical protein FLJ20825
1271	CTCATCTGCTG	10,00			syndecan 1
	AGGACATAATT	1,00		Hs.93454	ESTs
1273	GAGCTACACCA	1,00	0,34	Hs.82171	Human clone 191B7 placenta expressed mRNA from chrom
1274	TTGCTACTAAA	1,00	0,34	Hs.7790	ESTs
1275	CTTAGTGTTTT	1,00		Hs.7720	dynein, cytoplasmic, heavy polypeptide 1
1276	CTGGTCCTGGA	1,00	0,34	Hs.76476	cathepsin H
	GGTGGCAGTTG	1,00		Hs.75794	endothelial differentiation, lysophosphatidic acid G
1278	CTGATATAGAC	1,00	0.34		nuclear receptor coactivator 1
	TGGAAATCATT	1,00		Hs.5028	DKFZP564O0423 protein
	ACTITGAAAGG	1,00			hypothetical protein FLJ10793
	AAAGGCACTGA	1,00		Hs.3994	ESTs
	GTTCTCTTTTT	1,00		Hs.3843	dual specificity phosphatase 7
	AGCGCAGCTGT	1,00		Hs.34771	ESTs
	ATTGTGAAGAG	1,00			alpha2,3-sialyltransferase
	GCCTTCGGAAA	1,00		Hs.33104	Homo sapiens mRNA; cDNA
		.	·		DKFZp434H2121 (from clone DK
1286	GCTATTTTGAT	1,00	0,34	Hs.32250	ESTs
	TATCTCTTAAA	1,00		Hs.286228	ESTs
1288	TGTGATTTTA	1,00		Hs.286163	
1289	TATITCAGATT	1,00		Hs.285585	
	ATGATTTTGAG	1,00			putative selenocysteine lyase
1291	AATATTCATAT	1,00			Homo sapiens clone 25038 mRNA
		<u> </u>			sequence
	AGACCCCATTT	1,00		Hs.279297	
	CACCCATAGTC	1,00		Hs.278018	<del> </del>
	GTCTTGCTGCA	1,00			KIAA1171 protein
	CGGCCCATCTG	1,00		Hs.26290	
	CAATCTGATGC	1,00			hypothetical protein FLJ10261
1297	ATGTTGGGTGT	1,00	0,34	Hs.260855	Homo sapiens mRNA; cDNA DKFZp761G2311 (from clone DK
1208	ACTCTGTCTCC	1,00	U 34	Hs.259339	
	GGAATACAGAA	1,00			ESTs, Highly similar to vacuolar protein
1233	TOWN INCUGAN	1,00	0,34	JI 19.200020	ILO 19, mighty stitular to vacuolar protein

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					sorting hom
	ACTGGGCAAGC	1,00			hypothetical protein
	GGTCAGAAATT	1,00			metal-regulatory transcription factor 1
	AACCGAAGGGA	1,00		Hs.20596	
	GGGATAGAGAC	1,00			hypothetical protein FLJ20507
1304	TTTCAGTTAGT	1,00		Hs.196284	
1305	GCTATTGATGT	1,00		Hs.193398	
1306	AAGAGGAGGCC	1,00			hypothetical protein FLJ10210
1307	TAATACACTAA	1,00	0,34	Hs.183475	Homo sapiens clone 25061 mRNA
					sequence
1308	GCAGATGCTTT	1,00	0,34	Hs.180799	ESTs, Moderately similar to
					ALU5_HUMAN ALU SUBFAMILY
1309	TCACAAGGCTG	1,00		Hs.17998	
1310	TGCGAGCTGGG	1,00			collagen, type I, alpha 2
1311	GGATTTGCTGC	1,00	0,34	Hs.177956	Homo sapiens mRNA; cDNA
					DKFZp434C0926 (from clone DK
1312	CACGCACACAC	1,00			KIAA0914 gene product
1313	ATGTGGACTGA	1,00			KIAA0033 protein
1314	GGAGGCAGAGC	1,00	0,34		Human clone Z'3-1 placenta expressed
					mRNA from chrom
1315	GCTTTCTGTAA	1,00	0,34	Hs.172674	nuclear factor of activated T-cells,
					cytoplasmic 3
<del></del>	TGATTATTTAC	1,00		Hs.16930	
$\overline{}$	ACATCTGCCTG	1,00			hypothetical protein FLJ20159
1318	CTTAGTTTTAA	1,00			hypothetical protein FLJ20159
1319	AGGAAGAGTCA	1,00		Hs.154655	
1320	CAATGCAGAGG	1,00			malonyl-CoA decarboxylase
	TAATTCTTGTA	1,00		Hs.146123	
1322	CAAGGGCCCAC	1,00	0,34	Hs.14587	ESTs, Weakly similar to AF151859_1
<u></u>					CGI-101 protein [
	TTTTGAAGAAA	1,00		Hs.144465	
1324	TGTCTCCGTCT	1,00	0,34	Hs.135150	lung type-I cell membrane-associated
					glycoprotein
	TTTTCTTCAGG	1,00			chromosome 2 open reading frame 2
	TTCCTCCCTCT	1,00		Hs.125384	
1327	TCTGCCTTTCT	1,00	0,34	Hs.125019	ESTs, Highly similar to KIAA0886 protein
					[H.sapiens]
	ATAACTGTCAG	1,00			STE20-like kinase
-	TTGCAGTTTTT	1,00			CGI-43 protein
	TATTTAAAAAA	1,00		Hs.117304	
1331	GGCTCAGGGGC	1,00	0,34	Hs.116489	ESTs, Weakly similar to GCP170
					[H.sapiens]
	TTTATTGAAAC	1,00			mutS (E. coli) homolog 5
	CTGGCTTAAAT	1,00			APG5 (autophagy 5, S. cerevisiae)-like
1334	GCTTTATGTGG	1,00	0,34	Hs.111460	ESTs, Weakly similar to Con1
					[H.sapiens]
	GCATACTTTAT	1,00		Hs.109370	
	AGAATACTGAG	1,00	0,34	Hs.106705	neuronal PAS domain protein 2
1337	ACCCAAAAAA	1,00	0,34	Hs.101840	major histocompatibility complex, class I-

	,	, , , , ,			
					like seque
1338	GGACATTAGGG	1,00	0,34	Hs.101265	(Manual assignment) MEMOREC NSM2 (CCA1) sphingomyeli
1339	AAAAATAAATT	1,00	0,34	Hs.100261	Homo sapiens mRNA; cDNA DKFZp564B222 (from clone DKF
1340	AGATACATAGC	5,00	1,53	Hs.84045	Homo sapiens cDNA FLJ20288 fis, clone HEP04414
1341	TGGATATCAGT	5,00	1.53	Hs.7327	claudin 1
	TTTTCCACTTT	5,00		Hs.6900	ring finger protein 13
	GTGGCTCAGGC	4,00		Hs.259047	
	CCTGTGATTCC	4,00		Hs.227961	
	GGCGACAGAGC	11,00			hypothetical protein FLJ20163
	сствтветсст	11,00			Homo sapiens cDNA FLJ20463 fis, clone KAT06143
1347	TGCCTGTGGTC	16,00	4,15		ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB
	GTAAAAAAGCC	3,00	0,99	Hs.98988	ESTs
1349	TGTGAACACAT	3,00	0,99	Hs.80645	interferon regulatory factor 1
1350	AAACGAAGTTG	3,00	0,99	Hs.78353	SFRS protein kinase 2
	TACATCAGTAA	3,00	0,99	Hs.65029	growth arrest-specific 1
	CCTGTAGGCCC	3,00		Hs.207938	
1353	GTGAGACCTCG	3,00	0,99	Hs.203206	ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
	TGCCACCACAC	9,00	2,51	Hs.239993	ESTs
1355	GCTGGATTTTG	2,00	0,69	Hs.82124	laminin, beta 1
1356	TCACTTTTTA	2,00	0,69		ESTs
	ATTATCCTCAG	2,00	0,69	Hs.7987	DKFZP434F162 protein
1358	GGATCCAATTT	2,00	0,69	Hs.61796	transcription factor AP-2 gamma (activating enhancer
1359	CCAATTGAAGA	2,00	0,69	Hs.40328	ESTs
1360	TTACTTTTGGT	2,00	0,69	Hs.285861	hypothetical protein FLJ10359
1361	GAGAGCTTTGC	2,00	0,69	Hs.275374	aldo-keto reductase family 1, member C1 (dihydrodiol
	TACCCCCAAAC	2,00	0,69	Hs.241926	ESTs
1363	GGGCAGACACT	2,00			ESTs, Weakly similar to dJ876B10.4 [H.sapiens]
	ATGGCGCACGC	2,00			ESTs, Moderately similar to unnamed protein product
	AGGTTGCCGAG	2,00			KIAA0809 protein
	CCACTGCACCC	20,00	5,00	Hs.6853	carbohydrate (N-acetylglucosamine 6-O) sulfotransfer
1367	ATGGTGGGGA	18,00	4,55		zinc finger protein homologous to Zfp-36 in mouse
1368	СССТСТСССТТ	8,00	2,24	Hs.85087	latent transforming growth factor beta binding prote
1369	TCACCAAAAA	6,00	1,75	Hs.84753	KIAA0246 protein
	GTGAAACCCCC	59,00		Hs.265865	
1371	CCTGCAATCCC	34,00		Hs.3280	caspase 6, apoptosis-related cysteine protease

1372	GTGAAGCCCCG	22,00	5.34	Hs.285592	Homo sapiens mRNA; cDNA
		,	-,		DKFZp564M113 (from clone DKF
1373	CCACTGTACTC	53,00	11,94	Hs.220261	ESTs, Moderately similar to
1000	 		44.55		ALU4_HUMAN ALU SUBFAMILY
1374	стветвевсе	50,00	11,28	Hs.136810	ESTs, Weakly similar to ALU1_HUMAN
4075	10717010071	0.00	0.00	74040	ALU SUBFAMILY J S
	AGTATGACCTA	3,00			cytochrome c oxidase subunit VIc
13/6	GTGACAGCCAC	3,00	0,96	Hs.74441	chromodomain helicase DNA binding protein 4
1377	GGGCTTTTGAG	3,00	0.06	Hs.29893	Homo sapiens mRNA full length insert
1377	OCCOTTTOAG	3,00	0,90	1 13.23033	cDNA clone EURO
1378	GTGAGACCCCT	3,00	0.96	Hs.269952	ESTs, Weakly similar to ALU1_HUMAN
		,,,,,	-,		ALU SUBFAMILY J S
1379	GTGGTGCACAT	3,00	0,96	Hs.269030	
_	CCTGTAGTCAC	3,00		Hs.268900	
1381	TGGTAACTGGC	3,00	0,96	Hs.108741	ESTs
1382	GTGGTATGTGC	5,00	1,47	Hs.277102	ESTs, Weakly similar to ALU1_HUMAN
					ALU SUBFAMILY J S
	GTAAGATTAGC	5,00		Hs.250705	
1384	GCGAAACCCCA	72,00	15,71	Hs.210682	ESTs, Weakly similar to ALU6_HUMAN
					ALU SUBFAMILY SP
1385	ATCGTGCCACT	20,00	4,81	Hs.7615	Homo sapiens mRNA; cDNA
1000	T0T0T44T000	11.55	0.05		DKFZp434N2030 (from clone DK
1386	TCTGTAATCCC	44,00	9,85	Hs.142	sulfotransferase family, cytosolic, 1A,
1207	TTACCCACCCT	44.00	2.05	LI <sub>2</sub> 74267	phenol-prefe
1307	TTAGCCAGGCT	11,00	2,00	Hs.71367	ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
1388	ACAAAACCCTG	4,00	1,21	Hs.268591	
	ATCTCGGCTCA	14,00		Hs.29809	Homo sapiens mRNA; cDNA
			, i		DKFZp434C185 (from clone DKF
1390	CCTGTAATGCC	13,00	3,26	Hs.7179	RAD1 (S. pombe) homolog
1391	CCACCGCACTC	22,00	5,18	Hs.222669	ESTs, Moderately similar to
					ALU4_HUMAN ALU SUBFAMILY
1392	GTGGTGTGC	38,00	8,50	Hs.27038	Homo sapiens mRNA; cDNA
					DKFZp434G2127 (from clone DK
1393	ATGAAACCCCA	27,00	6,22		ESTs, Weakly similar to ALUC_HUMAN
4004	0070740000		4.00		IIII ALU CLASS C
1394	CCTGTAGCCCC	7,00	1,92	Hs.277320	EST, Weakly similar to ALU6_HUMAN
1205	TTTTTAAAAAA	2.00	0.66	11- 77040	ALU SUBFAMILY SP S
_	TTTTTAAAAAA AAGGAGCAAGT	2,00		Hs.77840	annexin A4
1380	ANGUNGUNNU I	2,00	0,00	Hs.76688	carboxylesterase 1 (monocyte/macrophage serine ester
1397	ACTITITATG	2,00	0.66	Hs.697	cytochrome c-1
	ATTGAGCCACA	2,00		Hs.63290	2-hydroxyphytanoyl-CoA lyase
	ACCACAAAAAA	2,00		Hs.469	succinate dehydrogenase complex,
		_,55	-,		subunit A, flavopro
1400	ATCACAGCTCA	2,00	0,66	Hs.29590	ESTs
1401	TGGTTCCAGCT	2,00			ESTs, Weakly similar to alternatively
					spliced produc
1402	TGACTGGCTTT	2,00	0,66	Hs.274439	Homo sapiens cDNA FLJ11265 fis, clone

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1404 C 1405 A 1406 T	GTGGTGGACCC CTGCTGTACTC	2,00	0.66		PLACE1009158
1404 C 1405 A 1406 T	CTGCTGTACTC	2,00	0 66		
1405 A 1406 T			0,00	Hs.270901	ESTs /
1406 T		2,00	0,66	Hs.22826	tropomodulin 3 (ubiquitous)
	ATGATAATTAA	2,00	0,66	Hs.170142	ESTs
1407 0	TAAAATAAGGG	2,00	0,66	Hs.169487	Kreisler (mouse) maf-related leucine
1407			·		zipper homolog
	GAGAGAGAA	2,00	0,66	Hs.169391	
1408 T	TACCCTGAAAC	2,00	0,66	Hs.144018	ESTs
1409 A	ACTGCCCGCTG	12,00	3,03	Hs.81071	extracellular matrix protein 1
1410	GGTGAGCGTGT	5,00			EphB3
	STGGTGGATGC	5,00		Hs.277904	
	ACTGTGCCACT	5,00			hypothetical protein FLJ20159
	GTGGCAGGCAC	68,00			ESTs, Weakly similar to ALUC_HUMAN
		,	,		IIII ALU CLASS C
1414 C	CCCACTTGTAA	12,00	3,00	Hs.75922	brain protein I3
1415	3CCCTTTCTCT	13,00			endocytic receptor (macrophage
<b>i</b> 1					mannose receptor fami
1416 A	AGACCTCCTTC	6,00	1,67	Hs.281706	
1417 A	AGTGGTGGCTA	6,00	1,67	Hs.230	fibromodulin
1418 C	GACAGATGTA	3,00	0,93	Hs.75356	transcription factor 4
1419 G	STGGCGAGCAC	3,00		Hs.261831	
1420 A	ATGGTGTGTGC	3,00	0,93	Hs.193347	ESTs
1421 G	GACTGAGTCA	3,00	0,93	Hs.18387	transcription factor AP-2 alpha (activating
			·		enhancer
1422 G	STGAGTGCCCT	3,00	0,93	Hs.171872	DEAD/H (Asp-Glu-Ala-Asp/His) box
					polypeptide 8 (RNA
1423 G	GCGGAACCTCA	3,00	0,93	Hs.10700	hypothetical protein
1424 G	STGTGGGGGGC	41,00	8,95		junction plakoglobin
1425 G	STGAAACTCCA	32,00	7,12	Hs.140002	ESTs, Moderately similar to
					ALU7_HUMAN ALU SUBFAMILY
1426 C	CCACTACACTC	29,00	6,50		tumor necrosis factor (ligand)
					superfamily, member 1
1427 A	ACGGAAGTTTT	4,00	1,18	Hs.144974	ESTs, Highly similar to unnamed protein
					product [H.s
1428 G	STGAAACCCGT	13,00	3,18	Hs.278577	Homo sapiens mRNA; cDNA
					DKFZp564P073 (from clone DKF
	CAAACTGTGA	5,00		Hs.94881	ESTs
	CATCGAAAGTT	2,00			hypothetical protein
1431 A	AGTAATCATCA	2,00	0,64	Hs.75925	proteasome (prosome, macropain)
11000	1.71.170.070.0				inhibitor subunit 1
	ATAATCCTGG	2,00			ESTs
1433 G	STATTCCTAAA	2,00	0,64	Hs.5724	ESTs, Weakly similar to multi PDZ
44045	OTOGO A A COAT			11- /00//	domain protein MUP
	CTGGGAAGCAT	2,00			ESTs
	SATCAAAACTG	2,00			c21orf7 form A-D
1430	STGACAGGCGC	2,00	0,64	Hs.278879	ESTs, Moderately similar to
1427 6	CACCOTOOAA	2 00		11- 07000	ALUA_HUMAN !!!! ALU CLAS
	SCACCGTGGAA	2,00			transcriptional regulator protein
1438	TTAACTGTATT	2,00	0,64	Hs.180952	actin, beta

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1439	ATGTTAGAGAC	2,00	0,64	Hs.1592	CDC16 (cell division cycle 16, S. cerevisiae, homolo
1440	ATCCCATCACT	2,00	0.64	Hs.158126	
	ATCGCATCACT				
	GACTCTGGAGA	2,00		Hs.154567	
	AAACTGTTCAA	2,00			KIAA0256 gene product
-	ACCAACACGGG	2,00		Hs.109005	
	AAAGATGTATC	1,00			pyruvate dehydrogenase (lipoamide) beta
	AAAACAGCAAG	1,00			SON DNA binding protein
1446	TTTTCAGGTAA	1,00	·		protein phosphatase 2 (formerly 2A), catalytic subun
1447	TGCCTCCCAGC	1,00	0,32	Hs.90527	HSPC128 protein
1448	TAAGTGAACAT	1,00	0,32		collagen, type XV, alpha 1
1449	TTTTGCTCAGA	1,00	0,32	Hs.8102	ribosomal protein S20
1450	CATTCTCCCAG	1,00	0,32	Hs.79110	nucleolin
1451	GTTTCAGCACT	1,00	0,32	Hs.77502	methionine adenosyltransferase II, alpha
1452	GTAACTCTATG	1,00	0,32	Hs.7277	peroxisomal biogenesis factor 3
1453	GTTCTATTGTA	1,00	0,32	Hs.6909	DKFZP564G202 protein
1454	CTATATTGTAA	1,00	0,32	Hs.65919	ESTs
1455	GTGAAACATTG	1,00	0,32	Hs.6567	Homo sapiens mRNA; cDNA
					DKFZp434C136 (from clone DKF
1456	GTATTGAAGTT	1,00	0,32	Hs.6079	B cell RAG associated protein
1457	CTTTAGAAGCA	1,00			ESTs
	TGACTCCTCAA	1,00	0,32		mitogen-activated protein kinase kinase kinase
1459	CTTTTATGGAC	1,00	0,32	Hs.44833	ESTs
1460	TAAATCTACAA	1,00	0,32	Hs.44701	ESTs
1461	TTCCCAAATGA	1,00	0,32		Homo sapiens mRNA; cDNA DKFZp762O2215 (from clone DK
1462	GTAAGAGTTCT	1,00	0.32		KIAA1025 protein
	ATGCCATTGGA	1,00			ceroid-lipofuscinosis, neuronal 5
	AGGAAATGGAT	1,00			ESTs
	AACAAGCTGGG	1,00			RNA POLYMERASE I AND
		,,,,			TRANSCRIPT RELEASE FACTOR
1466	TGCACTTGAGA	1,00	0.32	Hs.29055	ESTs
	GCATTCTGGTT	1,00		Hs.286261	
	GTGGCCACCCT	1,00			KIAA0685 gene product
	AAGGTGGTTGT	1,00			trinucleotide repeat containing 15
	AGAACTACGTG	1,00			hypothetical protein PRO2221
	AATTTGGGAGA	1,00			PC326 protein
	GGGAAACCCCT	1,00		Hs.279408	
	CTTACTCTTGA	1,00		Hs.27342	
	ACCGTGCCACT	1,00		Hs.270667	
	AGGCTGGTTTA	1,00			cell cycle related kinase
	TCTTTTGGGAG	1,00		Hs.257312	
	ACTGATCTTGT	1,00			CTP synthase
	AAGTCTGTAGA	1,00		Hs.250863	
	ATGGGGAAAGA	1,00		Hs.24989	
	ACAGTGCCACT	1,00		Hs.246374	
. 700	, .3/ .C   COO/LO	1,00	0,02	1.10.2-10074	<u></u>

1481   TGTGGGGACAA	4 404	TCTCCCCAAA	4 00	0.20	11- 045047	FOT
			1,00	0,32	HS.245017	EST
1483   ATTCCCTCCAAA	1482	CAAGICICCAG	1,00	0,32	HS.241515	
1484 ATGTATAGGC	4400	TTOOOTOOAAA	4 00	0.00	11. 000707	
1485   GCTTGTTGCGG						
1486   TCTTCTTAATA						
1487   GCTCATTTCAG						
CDNA clone EURO   1488   AGTTCCAGACC   1,00   0,32   Hs.223935   EST   ESTs, Weakly similar to ORF YGL221c   Sc. cerevisiae   St. cerevisiae   Cerevisiae   St. cerevisiae   Cerevisiae   St. cerevisiae   St. cerevisiae   Cerevi						Genefinder [
1489   AGTCAGTGGGA	1487	GCTCATTTCAG	1,00	0,32	Hs.22870	
Scerevisiae	1488	AGTTCCAGACC	1,00	0,32		
1491   TACATTTGAAT	1489	AGTCAGTGGGA	1,00	0,32	•	[S.cerevisiae]
beta isofo	1490	TTTCCAATGGA	1,00	0,32	Hs.21756	translation factor sui1 homolog
1493   AGAAAGATGGA	1491	TACATTTGAAT	1,00	0,32	Hs.21537	
1493   AGAAAGATGGA	1492	TTTTCTGTATT	1,00	0,32	Hs.21356	hypothetical protein DKFZp762K2015
1494 TTTACCTTTGG         1,00         0,32 Hs.21108         ESTs           1495 TACGATATTCA         1,00         0,32 Hs.207776         aspartylglucosaminidase           1496 GCACTGGGGCA         1,00         0,32 Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1497 GCTGCTAGAAA         1,00         0,32 Hs.197751         KIAA0666 protein           1498 GGGTAGAGAGT         1,00         0,32 Hs.194478         hypothetical protein FLJ10788           1499 ATCGGCTCCCA         1,00         0,32 Hs.194478         Homo sapiens mRNA, cDNA DKFZp434O1572 (from clone DK           1501 GAAGAAAGACT         1,00         0,32 Hs.172506         myosin VB           1502 TGCCTGACAGG         1,00         0,32 Hs.169160 ESTs           1503 TTCCTGTAATC         1,00         0,32 Hs.169160 ESTs           1505 ATAAAGCCGAA         1,00         0,32 Hs.169460 proteasome (prosome, macropain) subunit, alpha type,           1505 ATAAAGCGGAA         1,00         0,32 Hs.159471 ZAP3 protein           1506 AGGCTGAGGCG         1,00         0,32 Hs.154244 deiodinase, iodothyronine, type II           1508 TTCATAAAAAA         1,00         0,32 Hs.154257 matrix metalloproteinase 19           1509 GTGGCTACAGT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAT         1,00 <td< td=""><td>1493</td><td>AGAAAGATGGA</td><td>1,00</td><td></td><td></td><td></td></td<>	1493	AGAAAGATGGA	1,00			
1495 TACGATATTCA         1,00         0,32 Hs.207776 aspartylglucosaminidase           1496 GCACTGGGGCA         1,00         0,32 Hs.206259 Homo sapiens mRNA for KIAA1190 protein, partial cds           1497 GCTGCTAGAAA         1,00         0,32 Hs.197751 KIAA0666 protein           1498 GGGTAGAGAGT         1,00         0,32 Hs.196437 hypothetical protein FLJ10788           1499 ATCGGCTCCCA         1,00         0,32 Hs.194478 Homo sapiens mRNA; cDNA DKFZp434O1572 (from clone DK           1500 ATAATGGAGTG         1,00         0,32 Hs.172506 myosin VB           1501 GAAGAAAGACT         1,00         0,32 Hs.169160 ESTs           1503 TTCCTGTAATC         1,00         0,32 Hs.169160 ESTs           1503 TTCCTGTAATC         1,00         0,32 Hs.16364 hypothetical protein FLJ10955           1505 ATAAAGCCGAA         1,00         0,32 Hs.156292 ESTs           1507 TAGTGCTCTCA         1,00         0,32 Hs.156292 ESTs           1508 TTCATAAAAAA         1,00         0,32 Hs.154057 matrix metalloproteinase 19           1509 GTGGCTACAGT         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTTCTT         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTTCTT         1,00         0,32 Hs.151251 ESTs           1512 ACTAAGTGCTA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2 <td< td=""><td>1494</td><td>TTTACCTTTGG</td><td></td><td>0,32</td><td>Hs.21108</td><td>ESTs</td></td<>	1494	TTTACCTTTGG		0,32	Hs.21108	ESTs
1496   GCACTGGGGCA	1495	TACGATATTCA				
1497 GCTGCTAGAAA         1,00         0,32 Hs.197751         KIAA0666 protein           1498 GGGTAGAGAGT         1,00         0,32 Hs.196437         hypothetical protein FLJ10788           1499 ATCGGCTCCCA         1,00         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from clone DK           1500 ATAATGGAGTG         1,00         0,32 Hs.17850         ESTs           1501 GAGAAAGACT         1,00         0,32 Hs.172506 myosin VB           1502 TGCCTGACAAG         1,00         0,32 Hs.169160 ESTs           1503 TTCCTGTAATC         1,00         0,32 Hs.167106 proteasome (prosome, macropain) subunit, alpha type,           1504 AAATATTAAAC         1,00         0,32 Hs.16364 hypothetical protein FLJ10955           1505 ATAAAGCCGAA         1,00         0,32 Hs.159471 ZAP3 protein           1506 AGGCTGAGGGG         1,00         0,32 Hs.154292 ESTs           1507 TAGTGCTCTCA         1,00         0,32 Hs.154057 matrix metalloproteinase 19           1508 TTCATAAAAAA         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTTCTT         1,00         0,32 Hs.151251 ESTs           1511 GATGGGGAAAT         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1512 ACTAAGTGCTA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT				0,32	Hs.206259	Homo sapiens mRNA for KIAA1190
1498   GGGTAGAGAGT	1497	GCTGCTAGAAA	1.00	0.32	He 197751	
1499   ATCGGCTCCCA						
1500   ATAATGGAGTG						Homo sapiens mRNA; cDNA
1501   GAAGAAAGACT   1,00   0,32   Hs.172506   myosin VB   1502   TGCCTGACAAG   1,00   0,32   Hs.169160   ESTs   1503   TTCCTGTAATC   1,00   0,32   Hs.167106   proteasome (prosome, macropain)   subunit, alpha type,   1504   AAATATTAAAC   1,00   0,32   Hs.159471   ZAP3 protein   TAAAGCCGAA   1,00   0,32   Hs.159471   ZAP3 protein   ZAP3 protein   1506   AGGCTGAGGCG   1,00   0,32   Hs.156292   ESTs   1507   TAGTGCTCTCA   1,00   0,32   Hs.154424   deiodinase, iodothyronine, type   II   1508   TTCATAAAAAA   1,00   0,32   Hs.154057   matrix metalloproteinase   19   1509   GTGGCTACAGT   1,00   0,32   Hs.151251   ESTs   1510   TAATCTTTCTT   1,00   0,32   Hs.151236   highly charged protein   eukaryotic translation initiation factor 2C,   1   1512   ACTAAGTGCTA   1,00   0,32   Hs.19488   cystein-rich hydrophobic domain 2   1514   AGTGTGGGACT   1,00   0,32   Hs.119488   cystein-rich hydrophobic domain 2   1514   AGTGTGGGACT   1,00   0,32   Hs.119488   cystein-rich hydrophobic domain 2   1515   ATAGTTTAGCA   1,00   0,32   Hs.112157   ESTs   1516   CACCGAGACCA   1,00   0,32   Hs.107169   insulin-like growth factor binding protein   5   1517   AAATGACAATA   1,00   0,32   Hs.104904   ESTs   1518   TGACCAGGCGC   1,00   0,32   Hs.1019   parathyroid hormone receptor 1	4500	ATA ATOO A OTO	4.00	0.00	11- 47050	
1502   TGCCTGACAAG						
1503         TTCCTGTAATC         1,00         0,32         Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1504         AAATATTAAAC         1,00         0,32         Hs.16364         hypothetical protein FLJ10955           1505         ATAAAGCCGAA         1,00         0,32         Hs.159471         ZAP3 protein           1506         AGGCTGAGGCG         1,00         0,32         Hs.156292         ESTs           1507         TAGTGCTCTCA         1,00         0,32         Hs.154424         deiodinase, iodothyronine, type II           1508         TTCATAAAAAA         1,00         0,32         Hs.154057         matrix metalloproteinase 19           1509         GTGGCTACAGGT         1,00         0,32         Hs.151236         highly charged protein           1510         TAATCTITCTT         1,00         0,32         Hs.14520         eukaryotic translation initiation factor 2C, 1           1511         GATGAGGGAAAT         1,00         0,32         Hs.112430         l-mfa domain-containing protein           1513         CGGTTATTTAA         1,00         0,32         Hs.119488         cystein-rich hydrophobic domain 2           1514         AGTGTGGGACT         1,00         0,32         Hs.112157         ESTs						
Subunit, alpha type,   1504   AAATATTAAAC						
1505 ATAAAGCCGAA         1,00         0,32 Hs.159471 ZAP3 protein           1506 AGGCTGAGGCG         1,00         0,32 Hs.156292 ESTs           1507 TAGTGCTCTCA         1,00         0,32 Hs.154424 deiodinase, iodothyronine, type II           1508 TTCATAAAAAA         1,00         0,32 Hs.154057 matrix metalloproteinase 19           1509 GTGGCTACAGT         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTCTT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAAT         1,00         0,32 Hs.14520 eukaryotic translation initiation factor 2C, 1           1512 ACTAAGTGCTA         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1	<u> </u>					subunit, alpha type,
1506         AGGCTGAGGCG         1,00         0,32         Hs.156292         ESTs           1507         TAGTGCTCTCA         1,00         0,32         Hs.154424         deiodinase, iodothyronine, type II           1508         TTCATAAAAAA         1,00         0,32         Hs.154057         matrix metalloproteinase 19           1509         GTGGCTACAGT         1,00         0,32         Hs.151251         ESTs           1510         TAATCTTTCTT         1,00         0,32         Hs.151236 highly charged protein           1511         GATGGGGAAAT         1,00         0,32         Hs.14520         eukaryotic translation initiation factor 2C, 1           1512         ACTAAGTGCTA         1,00         0,32         Hs.132739         I-mfa domain-containing protein           1513         CGGTTATTTAA         1,00         0,32         Hs.119488         cystein-rich hydrophobic domain 2           1514         AGTGTGGGACT         1,00         0,32         Hs.118821         CGI-62 protein           1515         ATAGTTTAGCA         1,00         0,32         Hs.107169         insulin-like growth factor binding protein           1517         AAATGACAATA         1,00         0,32         Hs.104904         ESTs           1518         TGACCAGGC			1,00			
1507 TAGTGCTCTCA         1,00         0,32 Hs.154424 deiodinase, iodothyronine, type II           1508 TTCATAAAAAA         1,00         0,32 Hs.154057 matrix metalloproteinase 19           1509 GTGGCTACAGT         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTCTT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAAT         1,00         0,32 Hs.14520 eukaryotic translation initiation factor 2C, 1           1512 ACTAAGTGCTA         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1						
1508 TTCATAAAAAA         1,00         0,32 Hs.154057 matrix metalloproteinase 19           1509 GTGGCTACAGT         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTTCTT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAAT         1,00         0,32 Hs.14520 eukaryotic translation initiation factor 2C, 1           1512 ACTAAGTGCTA         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1						
1509 GTGGCTACAGT         1,00         0,32 Hs.151251 ESTs           1510 TAATCTTCTT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAAT         1,00         0,32 Hs.14520 eukaryotic translation initiation factor 2C, 1           1512 ACTAAGTGCTA         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1						
1510 TAATCTTCTT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAAT         1,00         0,32 Hs.14520 eukaryotic translation initiation factor 2C, 1           1512 ACTAAGTGCTA         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1						
1510 TAATCTTTCTT         1,00         0,32 Hs.151236 highly charged protein           1511 GATGGGGAAAT         1,00         0,32 Hs.14520 eukaryotic translation initiation factor 2C, 1           1512 ACTAAGTGCTA         1,00         0,32 Hs.132739 I-mfa domain-containing protein           1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1						
1511         GATGGGGAAAT         1,00         0,32         Hs.14520         eukaryotic translation initiation factor 2C, 1           1512         ACTAAGTGCTA         1,00         0,32         Hs.132739         I-mfa domain-containing protein           1513         CGGTTATTTAA         1,00         0,32         Hs.119488         cystein-rich hydrophobic domain 2           1514         AGTGTGGGACT         1,00         0,32         Hs.118821         CGI-62 protein           1515         ATAGTTTAGCA         1,00         0,32         Hs.107169         insulin-like growth factor binding protein           1516         CACCGAGACCA         1,00         0,32         Hs.107169         insulin-like growth factor binding protein           1517         AAATGACAATA         1,00         0,32         Hs.104904         ESTs           1518         TGACCAGGCGC         1,00         0,32         Hs.1019         parathyroid hormone receptor 1			1,00	0,32		
1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1	1511	GATGGGGAAAT	1,00	0,32	Hs.14520	eukaryotic translation initiation factor 2C,
1513 CGGTTATTTAA         1,00         0,32 Hs.119488 cystein-rich hydrophobic domain 2           1514 AGTGTGGGACT         1,00         0,32 Hs.118821 CGI-62 protein           1515 ATAGTTTAGCA         1,00         0,32 Hs.112157 ESTs           1516 CACCGAGACCA         1,00         0,32 Hs.107169 insulin-like growth factor binding protein           1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1	1512	ACTAAGTGCTA	1,00	0,32	Hs:132739	I-mfa domain-containing protein
1514         AGTGTGGGACT         1,00         0,32         Hs.118821         CGI-62 protein           1515         ATAGTTTAGCA         1,00         0,32         Hs.112157         ESTs           1516         CACCGAGACCA         1,00         0,32         Hs.107169 insulin-like growth factor binding protein           1517         AAATGACAATA         1,00         0,32         Hs.104904         ESTs           1518         TGACCAGGCGC         1,00         0,32         Hs.1019         parathyroid hormone receptor 1	1513	CGGTTATTTAA				
1515 ATAGTTTAGCA       1,00       0,32 Hs.112157 ESTs         1516 CACCGAGACCA       1,00       0,32 Hs.107169 insulin-like growth factor binding protein         1517 AAATGACAATA       1,00       0,32 Hs.104904 ESTs         1518 TGACCAGGCGC       1,00       0,32 Hs.1019 parathyroid hormone receptor 1	1514	AGTGTGGGACT				
1516 CACCGAGACCA 1,00 0,32 Hs.107169 insulin-like growth factor binding protein 5  1517 AAATGACAATA 1,00 0,32 Hs.104904 ESTs  1518 TGACCAGGCGC 1,00 0,32 Hs.1019 parathyroid hormone receptor 1						
1517 AAATGACAATA         1,00         0,32 Hs.104904 ESTs           1518 TGACCAGGCGC         1,00         0,32 Hs.1019 parathyroid hormone receptor 1						insulin-like growth factor binding protein
1518 TGACCAGGCGC 1,00 0,32 Hs.1019 parathyroid hormone receptor 1	1517	AAATGACAATA	1,00	0.32	Hs.104904	
						<del> </del>

	T .	r			II, DR alpha
1520	GTGGCGGGAGC	6,00	1.62	Hs.68257	general transcription factor IIF,
1320	GIGGCGGGAGC	0,00	1,02	105.00257	polypeptide 1 (74k
1521	CACAAACCCCC	40.00	9 14	Un 5496	clone FLB5214
_	GAGAAACCCCG	40,00		Hs.5486	
1522	AAATGCGAACA	4,00	1,10	Hs.5672	ESTs, Weakly similar to Similarity to
1522	ATCCACCTCCC	2 00	0.04	Un 224656	Yeast D-lactat
	ATCCACCTGCC	3,00		Hs.231656	<del></del>
	GCCAGCTGACA	3,00		Hs.118913	
	AGAAAGAATCT	3,00			small membrane protein 1
	TGCCCCTTGCC	3,00			secreted frizzled-related protein 4
	GAGAAACCCTG	35,00			ESTs, Weakly similar to ALUC_HUMAN IIII ALU CLASS C
	TTGCTGACTTT	36,00	7,54	Hs.108885	collagen, type VI, alpha 1
1529	AGGCTGAGGCA	11,00	2,64	Hs.17834	downstream neighbor of SON
1530	TCACTGCATTC	5,00	1,37	Hs.235587	EST
1531	CTTAAAAAAAA	5,00			hypothetical protein EDAG-1
1532	GTGAAAACCTG	9,00			ESTs, Moderately similar to
					ALU5_HUMAN ALU SUBFAMILY
1533	AGCCACCGTGC	31,00			DKFZP434D146 protein
1534	GTGGCACATAC	6,00	1,57	Hs.205353	ectonucleoside triphosphate
					diphosphohydrolase 1
1535	CCATTGTACTC	15,00	3,39	Hs.108740	DKFZP586A0522 protein
1536	TGCCTGTAATC	51,00	10,11	Hs.167135	Homo sapiens cDNA FLJ10728 fis, clone NT2RP3001236
1537	TTAGCCAGGAT	19,00	4,15	Hs.211457	EST
1538	TAGGGAATGAA	3,00			ring finger protein 15
1539	AAAGCATTTCT	3,00			ESTs, Moderately similar to WAP four- disulfide core
1540	ATGACCCGCAG	3,00	4,35	Hs.286254	ESTs, Weakly similar to AF170723_1
15/1	ATTITIONO	2.00	4.05	11- 070004	protein kinase ST
	ATTITITICAG	3,00		Hs.278004	
	GCAAGCCATTT	3,00			dual oxidase 1
	GATTTTTTTT	3,00		Hs.227913	
	TCTCTTGGGGT	3,00			hypothetical protein FLJ11036
	TGTGTGTAACA	3,00		Hs.156457	
	CCTTTGTCTTT	3,00		Hs.99654	protein-O-mannosyltransferase 1
	AATTGTAGTTA	3,00	_		RAP2A, member of RAS oncogene family
1548	AGCCACTGTAC	3,00	0,88	Hs.55346	ESTs, Weakly similar to Z141_HUMAN ZINC FINGER PROTE
1549	CAAACTCAAAA	3,00	0,88		hypothetical protein PRO1741
1550	GAGCACTTGGG	3,00			peptidylprolyl isomerase A (cyclophilin A)
1551	AAGTTTTTAGT	3,00		Hs.149917	
	TTTGAGGATTG	3,00		Hs.147916	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 3
1553	TAGTTGGAACT	3,00	0,88	Hs.1119	nuclear receptor subfamily 4, group A, member 1
1554	TTGACCAGGCT	13,00	2 08	Hs.285080	
	CTTATTTGTTT	5,00			plastin 3 (T isoform)
	OTTAIN SITE	5,00	1,34	110.7114	piasuit 5 (1 isolottii)

1556	AGCTTCCAGCC	5,00	1,34	Hs.144974	ESTs, Highly similar to unnamed protein
	<u>.</u>				product [H.s
	CACCCCCTCGC	2,00			hypothetical protein DKFZp547O146
1558	TCTCCAGGACA	2,00	0,61	Hs.8025	Homo sapiens clone 23767 and 23782 mRNA sequences
1559	TATTTCAATTG	2,00	0.61	Hs.79507	KIAA0582 protein
	CAGGTTGAAGT	2,00			RalGDS-like gene; KIAA0959 protein
	TATGTTAATGT	2,00		Hs.7341	ESTs, Weakly similar to DUS8_HUMAN DUAL SPECIFICITY
1562	GACTGCTCTGG	2,00	0.61	Hs.36475	ESTs
	GAAGAGTGCTC	2,00			ESTs
1564	GAGCCAAAGAA	2,00			ESTs, Weakly similar to macrophage lectin 2 [H.sapie
1565	CTTGTAATCTC	2,00	0,61	Hs.278002	EST
1566	TTACAATCACA	2,00		Hs.21276	
1567	GTGAAATCCAG	2,00	0,61	Hs.183275	ESTs
1568	CCTGTAATACC	9,00	2,17	Hs.92254	hypothetical protein FLJ20163
1569	AAAAGCAGAAA	4,00	1,11	Hs.84728	Kruppel-like factor 5 (intestinal)
1570	TTTGGGCCTAA	18,00	3,88	Hs.230320	EST
1571	CCTGTGGTCCC	52,00	10,04	Hs.249373	Homo sapiens clone FLB2543
1572	TAGCTCCCTTG	5,00			myeloid/lymphoid or mixed-lineage leukemia (trithora
1573	GTGAGACCCTG	22,00	4,58	Hs.135756	polymerase (DNA-directed) kappa
	GCGAAACCCCG	65,00			hypothetical protein
1575	CCACTGCATTC	38,00			ESTs, Weakly similar to ALUC_HUMAN IIII ALU CLASS C
1576	GGGATCGCCCC	6,00	1,52	Hs.12865	p47
	TTCCAAGGCAG	9,00	_		topoisomerase (DNA) I
	ATGGTGGGTGC	9,00			ESTs, Weakly similar to ubiquitous TPR motif, Y isof
1579	ATCTTGGCTCA	7,00	1,72	Hs.86958	interferon (alpha, beta and omega) receptor 2
1580	GTGAAACACCG	10,00	2,32	Hs.207766	EST
1581	CGTTCATTCAT	3,00	0,86	Hs.6139	synaptogyrin 1
1582	CCATAATGTTG	3,00	0,86	Hs.39957	pleckstrin 2 (mouse) homolog
1583	CTCTACGCATT	3,00	0,86	Hs.278573	H-2K binding factor-2
1584	ATGCAGAGGTG	3,00	0,86	Hs.210706	ESTs, Weakly similar to AF211175_1 unknown [H.sapien
1585	GCCAACAGCAT	3,00	0,86	Hs.155606	paired mesoderm homeo box 1
	CCTGTAATCAC	8,00		Hs.266136	
	GGATATGTGGT	21,00		Hs.738	early growth response 1
	GCTCACACCTG	4,00			sialyltransferase 4A (beta-galactosidase alpha-2,3-s
1589	CATACAGAAAA	4,00	1,09	Hs.3107	CD97 antigen
$\overline{}$	TATCCCAGAAC	27,00			crystallin, beta B2
	CGCCTGTAATC	33,00		Hs.235083	
	GTGGCACACAC	49,00			hypothetical protein
	ATCATACCACT	6,00			EST, Weakly similar to ALUC_HUMAN

1504	CTCCCTCCAC	10.00	2.04	Hs.228163	ICOT .
	GTGGCGTGCAC	19,00			
1595	TTGCCCAGGCT	50,00	9,38	HS.50U2/	Homo sapiens mRNA; cDNA DKFZp586J1717 (from clone DK
1596	CCGGTAATCCC	7,00	1,70	Hs.272813	dual oxidase 1
	AGCCACTGTGC	24,00		Hs.180606	
1598	GTGGTGCACAC	48,00	8,93	Hs.272173	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
1599	CCCACTCTTTG	3,00	0,84	Hs.9414	KIAA1488 protein
	CAAAATCTTGA	3,00			fibrinogen, gamma polypeptide
1601	CCTGTAGACCC	3,00			inorganic pyrophosphatase
1602	TCCTGGTTATT	3,00		Hs.4084	KIAA1025 protein
1603	TGCTAGATTGG	3,00			myeloid/lymphoid or mixed-lineage leukemia (trithora
1604	AACCCGGGAGA	3,00	0,84	Hs.236241	
	TACTCGGTTGT	3,00		Hs.119394	
	GACGGGGTGGA	3,00			hypothetical protein
1607	ATTTGTGTGTA	2,00	0,59	Hs.94499	ESTs
1608	GTTCCAAGCAA	2,00	0,59	Hs.94011	ESTs, Weakly similar to MAGE-B4 [H.sapiens]
1609	CTATCTGTGGA	2,00	0,59	Hs.9176	ESTs
1610	GGCCCAGAGCC	2,00	0,59	Hs.91246	hypothetical protein DKFZp547O146
1611	TTGATGCCCAG	2,00	0,59		ESTs
1612	TATTGTTAAAA	2,00	0,59	Hs.7984	ESTs
1613	CAATCTTTCAA	2,00	0,59	Hs.78909	butyrate response factor 2 (EGF- response factor 2)
1614	CTTCCTTGTGT	2,00	0,59		KIAA1151 protein
1615	GACAGTGATAG	2,00	0,59	Hs.53913	hypothetical protein FLJ10252
1616	GGCCTCTGATG	2,00	0,59		PRO1575 protein
1617	GCCTCCCCCAC	2,00	0,59	Hs.40109	KIAA0872 protein
1618	GGAGCAGACGC	2,00	0,59	Hs.31718	Homo sapiens cDNA FLJ11034 fis, clone PLACE1004258
1619	CTGCCCTCTGC	2,00	0,59	Hs.27801	zinc finger protein 278
1620	GGCTCTTCTGG	2,00	0,59	Hs.27721	hypothetical protein FLJ20353
1621	TTGCAATAGGT	2,00	0,59		hypothetical protein FLJ11323
1622	TGATGATCATT	2,00			hypothetical protein FLJ10893
1623	CCCAAACGGTA	2,00	0,59	Hs.195453	ribosomal protein S27 (metallopanstimulin 1)
1624	TTGGCCAAGAT	2,00	0,59	Hs.19522	hypothetical protein PRO2849
1625	TTTACCTGTTG	2,00			dihydropyrimidinase-like 2
1626	TGTCAATGGGG	2,00			golgi autoantigen, golgin subfamily a, 2
	CTTCCGGGTAA	2,00			DKFZP586P1422 protein
1628	AAGGTTCTTCT	1,00	0,29	Hs.89695	insulin receptor
	TAATTTTAAAC	1,00	0,29	Hs.8861	ESTs
	TTAAATGCTCT	1,00	0,29	Hs.82501	similar to mouse Xrn1 / Dhm2 protein
1631	AGCTCCCAAGA	1,00	0,29	Hs.80475	polymerase (RNA) II (DNA directed) polypeptide J (13
1632	TGCTGCGGAAG	1,00	0,29	Hs.80306	Homo sapiens mRNA, clone:RES4-4
	GAGCAGTGCTG	1,00		Hs.7636	feline sarcoma (Snyder-Theilen) viral (v- fes)/Fujina

1634	CAAGGAAATGT	1,00	0.29	Hs.64840	ESTs
	GCAACACCGGA	1,00		Hs.63525	poly(rC)-binding protein 2
	GCCAAAGATGT	1,00		Hs.58636	squamous cell carcinoma antigen
1030	GCCAAAGATGT	1,00	0,29	118.30030	recognized by T cell
1637	GTTGCAGGCGC	1,00	0.20	Hs.5811	hypothetical protein FLJ20467
	TCTTTAAAAAA	1,00			ESTs
	TGTGCTTCTAG	1,00		Hs.38613	ESTs
	TATCAAAAAAA	1,00		Hs.32491	ESTs
	TAACTCCTAGT	1,00			ESTs
	TGATATTAAAG	1,00		Hs.30661	electron-transferring-flavoprotein
1042	IGAIAIIAAAG	1,001	0,29	HS.3000 I	dehydrogenase
16/3	AAACCAATTTT	1,00	0.20	Hs.30483	Homo sapiens mRNA; cDNA
1043	AAACCAATTTI	1,00	0,29	115.30403	DKFZp434O1311 (from clone DK
1644	AAGAACTCAGG	1,00	0.20	Hs.30250	v-maf musculoaponeurotic fibrosarcoma
1044	MUMCICAGG	1,00	0,29	ITS.30230	(avian) oncoge
16/5	CAGTCCCAAAA	1,00	0.20	He 20846	Human DNA sequence from clone
1045	CAGICCCAAAA	1,00	0,29	115.23040	717M23 on chromosome 2
1646	AAACCGGTCCC	1,00	0.20	He 285490	ESTs, Weakly similar to unnamed
1040	~~~	1,00	0,29	115.205490	protein product [H.s
1647	AACATTCCTAA	1,00	0.20	Hs.285429	
	TTAACATTTAT	1,00			hypothetical protein FLJ10504
	ATGGCGTGTGC	1,00		Hs.278880	
	GAGCTCTTCCT	1,00			cytochrome P450, subfamily IID
1030	GAGCICITCCI	1,00	0,29		(debrisoquine, sparte
1651	GCAAGACCTTG	1,00	0.20	Hs.273603	
	AGGTTAAGAGA	1,00		Hs.272046	
	CCCGTAATCTC	1,00			Homo sapiens mRNA; cDNA
1000	CCCGIAXICIC	1,00	0,28	HS.210002	DKFZp586D0924 (from clone DK
1654	GCGGCACGCAC	1,00	0.20	Hs.269867	
$\equiv$	GACTTCCAGCA	1,00	0,29	He 265168	ESTs, Moderately similar to
1000	GACTICCAGCA	1,00	0,29	115.205100	T10_MOUSE SER/THR-RICH P
1656	GCTGTTCTAAG	1,00	0.20	He 24422	regulatory factor X-associated protein
	GCACTGAGAAG	1,00			KIAA0185 protein
	GCGAGACCTTG	1,00		Hs.232157	
	TTGCATTCTCC	1,00			ESTs
	TGCTTGTAGTC	1,00			ESTs, Weakly similar to ALU1 HUMAN
1000	1001101A010	1,00	0,23		ALU SUBFAMILY J S
1661	AACGCAGCCTT	1,00	0.29		KIAA0229 protein
	TAGGAAACCTG	1,00			KIAA0547 gene product
	GGCTTTATTCT	1,00			Homo sapiens mRNA; cDNA
333	COUNTRIES	',55	0,29	10.200712	DKFZp434G0719 (from clone DK
1664	AATACTTCTCT	1,00	0.20	Hs.2003	T cell receptor beta locus
	GGTGACAGAGG	1,00		Hs.200235	
	AGAAAAAAAAT	1,00			ESTs, Weakly similar to ALU5_HUMAN
		',00	0,29	13.200037	ALU SUBFAMILY SC
1667	CATTGGCACTC	1,00	0.20	Hs 195614	splicing factor 3b, subunit 3, 130kD
	TTAATTAGCAA	1,00			Homo sapiens mRNA; cDNA
		',55	0,20	1. 10. 100000	DKFZp434K098 (from clone DKF
1669	CGCGTCAGAGC	1,00	0.29	Hs.182982	
	1000 OAGAGG	1,00	5,23	1. 10. 102002	(Boiling of

1670	AAAACTCGCCG	1,00	0.29	Hs 17969	KIAA0663 gene product
	TGTACTTTCCT	1,00			tubulin, beta polypeptide
	CAGACTGGGAG	1,00		Hs.1790	nuclear receptor subfamily 3, group C,
			·		member 2
1673	TTTCTGAAGGG	1,00	0,29	Hs.172910	ESTs, Highly similar to unnamed protein product [H.s
1674	TAAATAAGGAA	1,00	0,29	Hs.17235	ESTs
	GATGACAGAGT	1,00			kallikrein 3, (prostate specific antigen)
	GTTACCGAGTG	1,00			ESTs, Weakly similar to KIAA1317 protein [H.sapiens]
1677	GGTCAAATCAT	1,00	0,29	Hs.170162	KIAA1357 protein
-	GCTTCACTTCC	1,00		Hs.164303	
	GACAATACACC	1,00			hypothetical protein FLJ20159
	TTCCAAAAAAA	1,00			SWI/SNF related, matrix associated,
			·		actin dependent
1681	TGTGACCCCTC	1,00	0,29		hexokinase 3 (white cell)
1682	TTTGTGCCATT	1,00	0,29	Hs.155507	ESTs
1683	CCTTGCCCTAT	1,00	0,29	Hs.143746	ESTs
1684	AGAACAAATAA	1,00		Hs.135721	
1685	AGACTGTACTG	1,00	0,29	Hs.132348	ESTs, Weakly similar to diaphanous 1
		·	·		[H.sapiens]
1686	CCATCCCAGTG	1,00	0,29	Hs.127863	
1687	ACAGACTGTTA	1,00			tumor endothelial marker 7 precursor
1688	GGCGACCCATT	1,00			echinoderm microtubule-associated protein-like
1689	CAGGTCCCATT	1,00	0,29	Hs.11924	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
1690	ACAAAGAAAAG	1,00	0,29	Hs.118578	Homo sapiens cDNA FLJ20053 fis, clone COL00809
1691	ACTGATGCTCA	1,00	0.29	Hs.115467	
	TCCTCTTCAA	1,00	0,29	Hs.113987	lectin, galactoside-binding, soluble, 2 (galectin 2)
1693	AGGCAGCACTG	1,00	0,29	Hs.11112	ESTs
1694	CTACTGCACTC	22,00	4,42	Hs.185989	ESTs
1695	CCCAGCTAATT	29,00	5,59	Hs.251235	EST
	GTGGCACGCAC	22,00		Hs.228343	
1697	GCAAAATCCCA	9,00	2,04	Hs.268051	ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB
1698	GTGAAACCTTG	35,00	6,56	Hs.161554	hypothetical protein FLJ20159
	ACTGTAATCCC	10,00		Hs.127809	
	ATCGCACCACT	19,00		Hs.142569	
	GGGAAACCCCA	10,00			ESTs, Weakly similar to alternatively spliced produc
1702	GTGAAACCCCT	27,00	5.16	Hs.229364	
	CAGCAGCAAAA	5,00		Hs.285090	
	TAGAAGCCAAC	7,00		Hs.7905	SH3 and PX domain-containing protein SH3PX1
1705	пспппсп	4,00	1,04	Hs.250722	(Manual assignment) MUG, Myeloid- upregulated protein

					r
1706	CCTATAATCCT	4,00	1,04	Hs.158164 	ATP-binding cassette, sub-family B (MDR/TAP), member
1707	TAAACGTGGCA	3,00	0,82	Hs.284146	hypothetical protein DKFZp762N0610
	CAGAAGTCTTC	3,00	0.82		ESTs, Weakly similar to ALU7_HUMAN
		·	,		ALU SUBFAMILY SQ
1709	ACCAGCCAAAG	3,00	0,82	Hs.193090	ESTs, Highly similar to AF161437_1
		·			HSPC319 [H.sapien
1710	GAAATGGGGAA	3,00	0,82	Hs.173933	Homo sapiens mRNA for KIAA1439
l					protein, partial cds
1711	GTGTGGTATTC	3,00	0,82	Hs.172140	ESTs
1712	CCGAGTTTTTG	3,00	0,82	Hs.139709	ESTs
1713	GGCAAACTTTA	3,00	0,82	Hs.102497	paxillin
1714	CCACAGGGGAT	15,00	3,05	Hs.119571	collagen, type III, alpha 1 (Ehlers-Danios
4=4=	00707070				syndrome
1715	CCTGTGGTCTC	6,00	1,44	Hs.236504	EST, Weakly similar to ALU6_HUMAN ALU SUBFAMILY SP S
1716	CACCACCACGC	5,00	1,23	Hs.5862	hypothetical protein
1717	GTGAAACCCGG	12,00	2,51	Hs.229170	
1718	GTGAAACCCTA	29,00	5,33	Hs.152081	ESTs
1719	ATATGTATATT	4,00	1,02	Hs.75839	zinc finger protein 6 (CMPX1)
1720	GTGAAACCACA	4,00			hypothetical protein DKFZp547A023
	GGGATTAAAGC	4,00			melanoma adhesion molecule
1722	TACCTTTGCTA	4,00			nuclear receptor co-repressor 2
1723	AATGAATGAAA	2,00	0,57	Hs.8986	complement component 1, q
					subcomponent, beta polypep
1724	ATATTTCATTC	2,00	0,57	Hs.79402	polymerase (RNA) II (DNA directed)
1725	TGGAGGGGCAG	2,00	0.57	Hs.7306	polypeptide C (33 secreted frizzled-related protein 1
	TCGCGCAATAA	2,00			protease-activated receptor 3
-	CTCAAAATCAA	2,00			hypothetical protein FLJ20283
	TGTACATATGT	2,00			homolog of yeast CDH1/HCT1
	TGCAATGTTGT	2,00	0,57	He 171057	triple functional domain (PTPRF
11723	100/4/101101	2,00	0,57	118.17 1337	interacting)
1730	TCATTTTGTGA	2,00	0,57	Hs.154567	
1731	TAAAACGTGAA	2,00	0,57	Hs.12592	period (Drosophila) homolog 3
1732	TCATCTGCAAA	2,00			ESTs, Weakly similar to AF148856_2
					unknown [H.sapien
1733	GGCAAAATCTA	2,00	0,57	Hs.104627	Homo sapiens cDNA FLJ10158 fis, clone HEMBA1003463
1734	GACCTATCTCT	8,00	1.78	Hs.194431	
	GTGGTGCGTGC	29,00			X-ray repair complementing defective
					repair in Chine
	TCTTGAACAGC	3,00			protease-activated receptor 3
	GGCTTTGGTCT	3,00			ribosomal protein, large, P1
	CCACCACACCC	3,00	0,80	Hs.117582	CGI-43 protein
	ATGAAACCCCG	25,00			hypothetical protein FLJ11126
	CCACTGCACTG	31,00		Hs.193220	
1741	CCACTGCGCTC	12,00	2,45	Hs.260287	ESTs, Weakly similar to ALU7_HUMAN
L	L				ALU SUBFAMILY SQ

1742	GAGAAACCCCA	30,00	5,36	Hs.5719	chromosome condensation-related SMC-associated prote
1743	CCTGTAATCCT	63,00	10.50	Hs.165954	
	GACAGTCGGTG	4,00		Hs.8203	endomembrane protein emp70 precursor isolog
1745	TTTTCTCTGAA	4,00	1,00	Hs.75516	tyrosine kinase 2
	TTGGCTAGGCC	7,00	1,57	Hs.211539	eukaryotic translation initiation factor 2, subunit
1747	CCCTTGTCCGA	7,00			ESTs, Weakly similar to weak similarity to collagens
1748	AGCCCAGGAGT	9,00	1,91	Hs.274813	EST
1749	GTGGTGTGCAC	17,00	3,18	Hs.20126_	KIAA0317 gene product
	CCTGTGATCCT	5,00			potassium channel, subfamily K, member 6 (TWIK-2)
	CCTGTAAACCC	5,00	1,17	Hs.161554	hypothetical protein FLJ20159
	TCAATAAAACC	4,00			KIAA0916 protein
	ACGAAACCCCA	4,00			CGI-43 protein
1754	TGACCACCCTT	3,00	0,78	Hs.42390	nasopharyngeal carcinoma susceptibility protein
1755	CTCGAATAAAA	3,00	0,78	Hs.34871	KIAA0569 gene product
1756	CGACTGCACTC	3,00	0,78	Hs.182061	Novel human gene mapping to chomosome 22
1757	CAGAATAATGT	3,00	0,78	Hs.125031	choline/ethanolaminephosphotransferase
	CAGAAGGCCAC	2,00		Hs.8268	ESTs
1759	GAAAGAGCTCT	2,00		Hs.7337	hypothetical protein FLJ10936
	AAAATTATCTT	2,00	0,55	Hs.63657	hypothetical protein FLJ11005
1761	GGTGTCTGTGG	2,00	0,55	Hs.5889	ESTs, Weakly similar to AC004876_5 similar to predic
1762	AACTGAGAAGT	2,00	0,55	Hs.56406	ESTs, Highly similar to unnamed protein product [H.s
1763	TGAGTGGTTTG	2,00	0,55	Hs.29672	ESTs
1764	GAAGTTGCCTT	2,00	0,55	Hs.26777	KIAA0843 protein
1765	TTGTTAAGCCT	2,00	0,55	Hs.26243	Homo sapiens cDNA FLJ11177 fis, clone PLACE1007402
1766	TATCTCAGAAC	2,00	0,55	Hs.223142	ESTs
1767	GGTGAATTTTA	2,00	0,55	Hs.210866	EST
1768	TGAGCACATAA	2,00	0,55		suc1-associated neurotrophic factor target 2 (FGFR s
1769	GTGCGTGCCTG	2,00	0,55	Hs.182354	ESTs
1770	ATTATCCAGCG	2,00	0,55	Hs.182225	RNA binding motif protein 3
1771	TCTTCTTTCAG	2,00			Homo sapiens mRNA; cDNA DKFZp434E1515 (from clone DK
1772	CTCTCCAAACC	2,00	0,55	Hs.151242	complement component 1 inhibitor (angioedema, heredi
1773	CCATTGCTCTC	2,00	0,55	Hs.117582	CGI-43 protein
	AAGATCCTTGT	2,00			karyopherin (importin) beta 3
	GGAACTTGGCT	2,00		Hs.105613	
	AGTTTGTCACC	2,00		Hs.10130	
	TCCACAGTGGG	1,00			ESTs, Weakly similar to 154197

		, <sub>T</sub>			hypothetical protein
1770	TACCCCTCTCA	4.00	0.07	He 004	hypothetical protein
	TACCCCTCTCA	1,00		Hs.994	phospholipase C, beta 2
	AAAGATGTACA	1,00	•	Hs.95243	transcription elongation factor A (SII)-like
1780	ATTTATAATCC	1,00	0,27	Hs.914	major histocompatibility complex, class II, DP alpha
1781	ACTGTTTGTTT	1,00	0,27	Hs.814	major histocompatibility complex, class II, DP beta
1782	TCGATGTGGCG	1,00	0,27	Hs.81248	CUG triplet repeat, RNA-binding protein
1783	TTAAGATCTTC	1,00	0,27	Hs.79404	neuron-specific protein
1784	TTCTTGTCATA	1,00		Hs.79081	protein phosphatase 1, catalytic subunit, gamma isof
	AACAATTATCA	1,00		Hs.7845	Homo sapiens cDNA FLJ20820 fis, clone ADSE00490
	AGAAACACTCA	1,00		Hs.75782	general transcription factor IIIC, polypeptide 2 (be
	AGAAATAAAAA	1,00		Hs.74649	cytochrome c oxidase subunit VIc
1788	TAAGTGTGGTT	1,00	0,27	Hs.7327	claudin 1
1789	AGGAGAGAGCC	1,00	0,27	Hs.6932	Homo sapiens clone 23809 mRNA sequence
1790	TGGACAAGTCA	1,00	0,27	Hs.64988	ESTs
1791	TTAAACTGCTG	1,00		Hs.6232	KIAA0764 gene product
1792	AGACCTCACTG	1,00		Hs.49763	ESTs
1793	TATTTGTATTT	1,00		Hs.4764	KIAA0763 gene product
	TGGATAGATTC	1,00		Hs.45519	ESTs
	TTGATTGATTT	1,00		Hs.42927	Homo sapiens cDNA FLJ11298 fis, clone PLACE1009794
1796	ACTATATTGTG.	1,00	0.27	Hs.42532	ESTs
	GAGTCCGGCCT	1,00		Hs.4069	glucocorticoid modulatory element binding protein 1
1798	TATTTATTTT	1,00	0,27	Hs.39143	ESTs, Weakly similar to predicted using Genefinder [
1799	TGACATCCTGA	1,00	0,27	Hs.285056	ESTs
	CTGCAAGGACA	1,00			Homo sapiens HSPC295 mRNA, partial cds
1801	ATCCCCCAGAA	1,00	0,27	Hs.278386	ESTs
1802	CCACTGCGCTT	1,00	0,27	Hs.252836	EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SE
1803	TGCCAGACCCT	1,00	0,27	Hs.249721	
1804	GGTGTGCACCT	1,00			signal transduction protein (SH3 containing)
1805	CTTTATTTT	1,00	0,27	Hs.245710	heterogeneous nuclear ribonucleoprotein H1 (H)
1806	AGCGCTGGGGA	1,00	0,27	Hs.241471	
	CTAGGACCTGT	1,00	0,27	Hs.240112	KIAA0276 protein
1808	TAGTCCTAGCT	1,00		Hs.237372	
	ATTTAATTTA	1,00		Hs.235883	
1010	GGCAACAAAGT	1,00		Hs.233364	

1811	CTGTAAGGATC	1,00	0.27	Hs 227730	integrin, alpha 6
	AGTCTTCCAGT	1,00			cytochrome P450, 51 (lanosterol 14-
1012	AGIOTIOOAGI	1,00	0,21	10.220213	alpha-demethylase
1812	GCTCCCCCTCC	1,00	0 27	Hs.2157	Wiskott-Aldrich syndrome (ecezema-
1013	30100000100	1,00	0,21	10.2101	thrombocytopenia)
1914	AGTATTTATGA	1,00	0.27	Hs.203838	
	GCTAAACCCTG	1,00			ESTs, Moderately similar to
1010	GCTAAACCCTG	1,00	0,27	NS.202761	ALU5_HUMAN ALU SUBFAMILY
1016	CCAGCATTACC	1,00	0.27	Hs.20082	Homo sapiens zinc finger protein NY-
1010	CCAGCATTACC	1,00	0,27	П5.20002	REN-21 antigen m
1817	CCTGCAATCTC	1,00	0.27	Hs.197793	
	GAAAAATGCGC	1,00		Hs. 193398	
	GCCAGGGCTCA	1,00			ESTs, Moderately similar to MRP3
1019	GCCAGGGCTCA	1,00	0,21	18. 10 <i>19</i> 13	[H.sapiens]
1820	TAAAACTTACA	1,00	0.27	Hs.184075	
	ACCCTTTTTAT	1,00			ADP-ribosylation factor 4-like
	GTTTCAAACGA	1,00			ESTs, Weakly similar to S69890 mitogen
1022	GIIICAAACGA	1,00	0,27		inducible gen
1822	CAGTCTCAGTG	1,00	0.27		Homo sapiens mRNA; cDNA
1023	CAGICICAGIG	1,00	0,21	П5. 17 / 07	DKFZp761N07121 (from clone D
1924	AAAAATTCATC	1,00	0.27	He 170229	
				Hs.170328	
	TAAATAAACAA	1,00	0,27	HS. 10/55	MBIP protein
1826	CACCTCAAACA	1,00	0,27	HS.15/150	ESTs, Weakly similar to zinc finger
4007	AATOTOOTOOO	4 00	0.07	11- 455007	protein 106 [M.m
	AATGTCCTCGG	1,00			KIAA0645 gene product
1828	TGTACCCCGCT	1,00	0,27	Hs.155975	protein tyrosine phosphatase, receptor
1000	00007770700	4 00		11 455400	type, C-assoc
_	CGGGTTTGTGC	1,00	0,27	Hs.155482	hydroxyacyl glutathione hydrolase
	GTCCATCTTAA	1,00			ribosomal protein S28
	TTTATTTTTAG	1,00		Hs.152250	
1832	CCTAAAAAAAA	1,00	0,27	Hs.148907	Homo sapiens mRNA; cDNA
1000					DKFZp564G223 (from clone DKF
	TGATCGAGCTT	1,00		Hs.145867	
1834	GAAGATATTCC	1,00	0,27	Hs.14553	sterol O-acyltransferase (acyl-Coenzyme
1005	1=00100=00	1.55			A: cholester
	ATGGAGCTGCA	1,00	0,27	Hs.142779	ESTs
	ATAAGACCTTA	1,00	0,27	Hs.142296	jerky (mouse) homolog
	CAAGCCAAAAA	1,00			hypothetical protein FLJ10379
	CAAATGGCAAA	1,00		Hs.134292	
1839	GAAGTTTAAAT	1,00	0,27	Hs.132463	phosphoinositide-3-kinase, class 2, beta
					polypeptide
	GAAGGCAAGAT	1,00		Hs.1321	coagulation factor XII (Hageman factor)
	TTAGTTATGAC	1,00			transcription factor CA150
	CTGTATGTTTA	1,00		Hs.128777	
1843	ACCTGCATTCC	1,00	0,27	Hs.125034	Homo sapiens cDNA FLJ10733 fis, clone
					NT2RP3001392
1844	CCAGCTGCCTG	1,00	0,27	Hs.11782	ESTs
1845	AAAAGTGGTGT	1,00			CGI-43 protein
1846	TTTATCTGATA	1,00			CGI-43 protein

or mixed-lineage
oduct
-Ala-Asp/His) box
yc
RNA; cDNA
(from clone DK
NA; cDNA
from clone DKF
pen reading frame 50
ar to PMM2 HUMAN
OMUTAS -
ypeptide 9, non-
; small stress protein-
, a
-binding, soluble, 3
·
chromosome region,
ne 23632 mRNA
n FLJ20686
nate-5-phosphatase,
spindle assembly 1
se
NA; cDNA
(from clone DK
ma cell differentiation-
se
eotide exchange factor
ator, urokinase
similar to
U SUBFAMILY
rotease 11
in

					·
	GGGAAACAGGT	4,00		Hs.18368	DKFZP564B0769 protein
	ССТІТІТІТІ	2,00		Hs.9956	hypothetical protein FLJ20259
	GGGGCTTAGGA	2,00		Hs.89135	KIAA1528 protein
	CTAGACAGTAA	2,00			KIAA0669 gene product
1884	GTGTTCTGTGC	2,00	0,53	Hs.241567	RNA binding motif, single stranded
					interacting prote
1885	TGCTGTAAAGG	2,00	0,53	Hs.23856	Homo sapiens HSPC091 mRNA, partial
					cds
	GAGGAGTGGGT	2,00	0,53	Hs.206770	zinc finger protein 297
	GTAAGACCCTG	2,00		Hs.164177	
1888	GGCCGTTAGAA	2,00	0,53	Hs.135	methylmalonate-semialdehyde
					dehydrogenase
	AGGCTAAAAGC	2,00	0,53	Hs.113029	ribosomal protein S25
$\overline{}$	CTGTGTAATTT	2,00		Hs.109731	
	CTGAAGCGTGC	2,00			Human insulin-like growth factor binding protein 5 (
1892	AGAACCTTCAA	9,00	1,78	Hs.181244	major histocompatibility complex, class I, A
1893	TTCTGTGCTGG	16,00	2,84	Hs.1279	complement component 1, r
					subcomponent
	TTAGCTGAGTC	5,00			cytochrome b-561
1895	TTGGCCAGACT	9,00	1,76	Hs.91728	polymyositis/scleroderma autoantigen 1
					(75kD)
1896	TTTCATTGCCT	9,00	1,76	Hs.173159	transforming, acidic coiled-coil containing
100=					protein
1897	GTGGCCAGAGG	9,00	1,76	Hs.1420	fibroblast growth factor receptor 3
4000	A000770707A	0.00	1.00	11 11 17 17 17 17	(achondroplasia,
	ACCGTTCTGTA	6,00			CGI-43 protein
	TAACTCCAAAG	3,00		Hs.24743	hypothetical protein FLJ20171
	TGCCGTAAATG	3,00			v-erb-b2 avian erythroblastic leukemia viral oncogen
	TGAACTTTCCT	3,00	0,74		ESTs
1902	TAAAGATCCTC	3,00	0,74	Hs.100407	Homo sapiens mRNA; cDNA
					DKFZp564H2416 (from clone DK
1903	GTGGCTCACAC	55,00	8,11	Hs.138411	Homo sapiens mRNA; cDNA
					DKFZp586J1922 (from clone DK
	GTGGTACACAC	5,00		Hs.250419	
	GATCTCTTGGG	5,00			keratin 16 (focal non-epidermolytic palmoplantar ker
	GAGGAACTCAA	7,00	1,40	Hs.5008	CGI-87 protein
	AATAAAGCCTT	6,00	1,24	Hs.3314	selenoprotein P, plasma, 1
	TTTACAAGTTA	4,00			hypothetical protein DKFZp547O146
1909	AGGTCAAAAAA	4,00	0,91		actin related protein 2/3 complex, subunit 4 (20 kD)
1910	GAGCCCCCGTG	4,00	0,91	Hs.12908	CDC42-binding protein kinase beta
					(DMPK-like)
1911	GAGTAGCTGAG	3,00	0,72	Hs.260039	sarcospan (Kras oncogene-associated
					gene)
1912	GTGCTGCTCCA	2,00	0,52	Hs.7936	BAI1-associated protein 2

4040	0.00	0.00	0.50	75450	[
	GAGATTTGTTT	2,00			heat shock 70kD protein 2
	GAAGGGGTGCT	2,00			DKFZp434A0131 protein
	GCCACAGTACA	2,00			DKFZP586H2123 protein
	TAATTTTACT	2,00			hypothetical protein FLJ20624
1917	ACTGTTTGGCA	2,00	0,52	Hs.286110	translocase of inner mitochondrial
					membrane 9 (yeast
1918	TCTGGCTAATT	2,00		Hs.262198	
1919	GTGGAAACCCA	2,00	0,52	Hs.243818	ESTs, Moderately similar to
					ALU1_HUMAN ALU SUBFAMILY
1920	TTGCCCAAGCT	2,00	0,52	Hs.213469	EST
1921	AAAACAGTGGC	2,00	0,52	Hs.184109	ribosomal protein L37a
1922	TGATGTGATAG	2,00	0,52	Hs.181159	Homo sapiens mRNA; cDNA
					DKFZp434F0217 (from clone DK
1923	TTTGAACCCTT	2,00	0,52	Hs.16206	uncharacterized hypothalamus protein
					HT008
1924	CCTATAATAAA	2,00	0,52	Hs.13885	ESTs, Weakly similar to T09A5.6
1					[C.elegans]
1925	AGAATCACTTA	2,00	0,52	Hs.130815	ESTs
	TATTTTGCAAA	2,00			DKFZP564O123 protein
	CCTATAACCCC	1,00			ESTs
	TCCAACTACAC	1,00		Hs.94581	sulfotransferase family, cytosolic, 2B,
1		,,	-,		member 1
1929	TACCCAAAGAA	1,00	0.26	Hs.9436	ESTs, Weakly similar to NC5R_RAT
		,,,,,	-,		NADH-CYTOCHROME B5
1930	TGTTTGTAAAA	1,00	0.26	Hs.9271	KIAA1071 protein
	пппппс	1,00		Hs.90797	Homo sapiens clone 23620 mRNA
		.,	-,		sequence
1932	TATCTCTGCAA	1,00	0.26	Hs.82985	collagen, type V, alpha 2
	TTCTTCTGAAA	1,00		Hs.8087	NAG-5 protein
	CAGATGTTTAA	1,00		Hs.77631	glycine cleavage system protein H
1.55		.,	ا - ر		(aminomethyl carri
1935	TTTGTAATATT	1,00	0.26	Hs.75546	capping protein (actin filament) muscle
1.000		.,00	0,20	110000	Z-line, alph
1936	ACCCAGTTGTT	1,00	0.26	Hs.75410	heat shock 70kD protein 5 (glucose-
1.000		.,,55	0,20		regulated protein
1937	CTCATTGGTGG	1,00	0.26	Hs.6580	Homo sapiens clone 23718 mRNA
1		.,,,,	0,0		sequence
1938	TCTTCTCACAA	1,00	0.26	Hs.656	cell division cycle 25C
	CCTTTGTTCAA	1,00		Hs.6107	ESTs
	TTAGAGATTCC	1,00		Hs.5947	mel transforming oncogene (derived from
.5 15		.,55	5,20	. 10.0071	cell line NK
1941	TCCACACCAAA	1,00	0.26	Hs.53656	ESTs, Weakly similar to D29149 proline-
	. 50/10/100/07	.,00	0,20	5.55555	rich protein
1942	TGTAATGGTTT	1,00	0.26	Hs.4930	low density lipoprotein receptor-related
1572		1,00	0,20	13.4000	protein 4
1943	GTACTTACCTT	1,00	0.26	Hs.3454	ESTs, Weakly similar to KIAA0665
1070		1,00	0,20	10.0707	protein [H.sapiens]
1944	CTTAAATGGTT	1,00	0.26	Hs.29679	cofactor required for Sp1 transcriptional
1,244		1,00	0,20	1 13.23013	activation
L	L			L	activation

40.00	0=004400=04	4.00	0.00	11. 005000	It is a total and a containing AE
	CTCCAACCTGA	1,00			trinucleotide repeat containing 15
	TTAGGCTTTAG	1,00			hypothetical protein FLJ20392
	GAAGATGTACG	1,00		Hs.285077	
	TGCCACCATAC	1,00		Hs.284138	
	TGTCTGTAGTC	1,00		Hs.282837	
	TAAAGTCCATT	1,00			KIAA1117 protein
	CCAACTGACTT	1,00			KIAA0631 protein
1952	ATGGAATGCTA	1,00	0,26	Hs.268551	receptor-interacting serine-threonine
					kinase 3
1953	TTGAAACCTCG	1,00		Hs.267148	
1954	TATATCATATT	1,00	0,26	Hs.266914	hypothetical protein FLJ10355
1955	TGGCACGCTGC	1,00	0,26	Hs.250890	ESTs, Weakly similar to TOM1
		_ 1			[H.sapiens]
1956	ACAGAGTCTCA	1,00	0,26	Hs.249031	EST
1957	GAAATATTGAT	1,00	0,26	Hs.247043	type 1 tumor necrosis factor receptor
ļ ļ		•	·		shedding amino
1958	GTGAAACCTGA	1,00	0,26	Hs.242076	EST
1959	ATGTCAACCAA	1,00	0,26	Hs.241558	ariadne (Drosophila) homolog 2
	GAAAAGGGCAC	1,00			KIAA1105 protein
-	TGCAGTCTTTG	1,00		Hs.232111	
	CAGATTTCCAG	1,00			ESTs, Weakly similar to AF121081_1
		",""	-,		cAMP inducible 2
1963	AAAAGGAAACC	1,00	0.26		Homo sapiens mRNA; cDNA
1.000		.,	-,		DKFZp761K2024 (from clone DK
1964	ATTGTAAGCTT	1,00	0.26	Hs.210232	
	GTCTTAAAATA	1,00			DKFZP564A122 protein
	TCCGCAGGGAA	1,00			Human clone A9A2BRB5
1.000	7000071000711	. 7,00	0,20	710.70 7002	(CAC)n/(GTG)n repeat-containing
1967	GCTGGAGCTCA	1,00	0.26	Hs.181315	ESTs, Moderately similar to
	001001071	1,50	0,20		ALU4 HUMAN ALU SUBFAMILY
1968	TGTTGTAAATA	1,00	0.26	Hs.171501	ubiquitin specific protease 11
	GCTCCTACATT	1,00			dentatorubral-pallidoluysian atrophy
1.000	0010017107111	1,00	0,20	110.100.100	(atrophin-1)
1970	CAGGTGCCAAA	1,00	0.26	Hs 168350	KIAA0554 protein
	TGCTGCTGCCC	1,00		Hs.167046	
<b>—</b>	GCCTGGGCTGA	1,00			hypothetical protein FLJ20626
	GTATGAGGTGG	1,00		Hs.164464	
	GTAAACAGAAA	1,00			hypothetical protein FLJ20159
	TGAAATAAACT	1,00			methylmalonyl Coenzyme A mutase
	TTTTGTCAACA				ESTs, Weakly similar to tyrosine kinase
11976	ITTTGTCAACA	1,00	0,20	ITS. 154645	[H.sapiens]
4077	COACCTOTTOT	4.00	0.26	Un 452006	cytochrome P450, subfamily IIJ
11977	GCACGTGTTCT	1,00	0,26	RS. 152096	
1070	TTACTCCACAC	4 00	0.00	He 450000	(arachidonic acid epo
	TTAGTCCACAG	1,00			zinc finger protein 262
19/9	AAATTTCAAGC	1,00	0,26	ms. 1464U1	small inducible cytokine subfamily E,
4000	00040074077	4 00		11- 440004	member 1 (endo
1380	CCCAGCTACTT	1,00	0,26	<sub> </sub> HS.143961	ESTs, Moderately similar to
4004	A A A C C A A A	4 00	0.00	11- 400400	ALU1_HUMAN ALU SUBFAMILY
[198]	AAACCAGGAAA	1,00	0,26	<sub>[HS.139120</sub>	ribonuclease P (30kD)

4000	IAATETCAACAA	4 00	0.00	11- 440504	Eductor related protein security 2 pinns
<u></u>	AATTTCAAGAA	1,00			adaptor-related protein complex 2, sigma 1 subunit
1983	TACCCTAAAAT	1,00	0,26	Hs.117325	Homo sapiens cDNA FLJ11166 fis, clone PLACE1007242
1984	GGACTGTAGTG	1,00	0.26	Hs.11711	KIAA0329 gene product
	CTCCCGCCGGA	1,00			KIAA1020 protein
	TGAAGTGCCCT	1,00		Hs.106932	
	GAAGAGGCTGG	1,00		Hs.105962	
	ттесеттетте	9,00		Hs.814	major histocompatibility complex, class II, DP beta
1989	AACCCGGAAGG	8,00	1,54		butyrophilin, subfamily 3, member A2
	GTGGCGCGTGC	16,00	2,66	Hs.24135	hypothetical protein DKFZp761C241
1991	TACCCTAAAAC	103,00			KIAA0675 gene product
$\overline{}$	CCTGTGATCCC	40,00			cathepsin B
	CCGGCCCTACC	4,00			epithelial protein up-regulated in carcinoma, membra
1994	GCACGCGTAAC	4,00	0,89	Hs.169552	ESTs, Weakly similar to BRDT [H.sapiens]
	AACAAGGTGAG	3,00			ESTs, Highly similar to transcription elongation fac
	TGCGTCCCTCC	3,00			DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k
1997	GACCCTTTTGG	3,00			Homo sapiens mRNA; cDNA DKFZp434G1310 (from clone DK
	GATTTTTCTGA	3,00			RNA binding motif, single stranded interacting prote
	TGATTCTGTTT	3,00			collagen, type V, alpha 1
2000	CCCGGCTAATT	14,00	2,35	Hs.102926	ESTs
2001	GTGAAGCCCTG	16,00			ectodermal dysplasia 1, anhidrotic
2002	ACTGAAAGAAG	5,00			complement component 1, s subcomponent
	CTGAGAGCTGG	14,00			growth arrest-specific 6
	GTTCCAGCAGC	4,00	_		Homo sapiens clone 25116 mRNA sequence
	CCATTGCGCTC	4,00	0,88	Hs.204299	ESTs, Moderately similar to alternatively spliced pr Homo sapiens mRNA full length insert
	GGATGCGCAGG	4,00			cDNA clone EURO
	TTGGTGGAGGT	2,00			CD63 antigen (melanoma 1 antigen)
2008	AGAATTATGGG	2,00			PRO1073 protein
	TGTGGTGGCAC	2,00	0,50	Hs.46624	HSPC043 protein
2010	CAGTTCTTGAT	2,00	0,50	Hs.284217	serologically defined colon cancer antigen 33
2011	GCAAGACCCCG	2,00	0,50	Hs.262335	ESTs, Weakly similar to ALUC_HUMAN
2012	AAAACAAAACA	2,00	0,50	Hs.24734	oxysterol binding protein
	AAGACTGACAA	2,00			topoisomerase-related function protein 4
	TTCTCCTCTTT	2,00			hypothetical protein FLJ10357
	TCAATCAGTGA	2,00		Hs.127270	
			-,		L

2016 TGGGGTCCCCA   2,00   0,50 Hs.123661 ESTs   STs, Moderately similar to altern   spliced pr   2018  AATAAATTCCT   14,00   2,30  Hs.76307   neuroblastoma, suppression of   tumorigenicity 1   2019  GTGGCAGATGC   7,00   1,34 Hs.273539 ESTs, Weakly similar to ALU2_Ht   ALU SUBFAMILY SB   2021  GCGAAACTCCA   7,00   1,34 Hs.12860 KiAA1353 protein   2021  GCGAAACTCCA   7,00   1,34 Hs.12860 KiAA1353 protein   2022  GTGAAACTCCA   7,00   1,34 Hs.12860 KiAA1353 protein   2022  GTGAAACTCCA   7,00   1,34 Hs.12860 KiAA1353 protein   2023  GGCAGACACAT   3,00   0,69 Hs.323267   nuclear factor I/B   2024 AAAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAG	
Spliced pr   neuroblastoma, suppression of tumorigenicity 1	natively
2018   AATAAATTCCT	
tumorigenicity 1	
2019   GTGGCAGATGC   7,00	
ALU SUBFAMILY SB	UMAN
2020   CCTGTTATCCC	
2021   GCGAAACTCCA	
2022 GTGAAACCTCG         38,00         5,28 Hs.194408 KIAA1244 protein           2023 GGCAGACACAT         3,00         0,69 Hs.33287 nuclear factor I/B           2024 AAAGAGAGAG         3,00         0,69 Hs.32969 ESTs           2025 ACAAAGCATTT         40,00         5,46 Hs.103391 Human insulin-like growth factor by protein 5 (           2026 TTTAGTGACGT         7,00         1,32 Hs.7104 Kruppel-like factor 13           2027 AGCCACCACGC         17,00         2,63 Hs.60772 ESTs           2028 GCGAAACCTCG         8,00         1,46 Hs.210473 ESTs, Weakly similar to GELS_HI GELSOLIN PRECURSO           2029 GCAGTTGGATC         4,00         0,86 Hs.284932 Homo sapiens clone 24650 ubiqui hydrolase mRNA, p           2030 CTTGTGAAGTG         4,00         0,86 Hs.283681 ESTs           2031 GTGGCTCACGC         56,00         7,21 Hs.228230 EST           2032 CAGCTATTTCA         10,00         1,69 Hs.153179 fatty acid binding protein 5 (psoria associated)           2033 CACACACACAC         4,00         0,84 Hs.256696 EST, Weakly similar to ALU1_HU LU LU SUBFAMILY J SE           2035 CAAGGGTGACA         4,00         0,84 Hs.276096 EST, Weakly similar to ALU1_HU LU LU SUBFAMILY J SE           2036 GCAGCACTTAT         2,00         0,48 Hs.76144 muskelin 1, intracellular mediator containing kelch           2037 CCTGACCTCAA         2,00         0,48 Hs.76144 muskelin 1, intracellular	
2023   GCAGACACAT   3,00   0,69   Hs.33287   nuclear factor I/B	
2024   AAAGAGAAGAG   3,00	
2025 ACAAAGCATTT         40,00         5,46 Hs.103391 Human insulin-like growth factor be protein 5 (           2026 TTTAGTGACGT         7,00         1,32 Hs.7104 Kruppel-like factor 13           2027 AGCCACACGC         17,00         2,63 Hs.60772 ESTs           2028 GCGAAACCTCG         8,00         1,46 Hs.210473 ESTs, Weakly similar to GELS_HI GELSOLIN PRECURSO           2029 GCAGTTGGATC         4,00         0,86 Hs.284932 Homo sapiens clone 24650 ubiqui hydrolase mRNA, p           2030 CTTGTGAAGTG         4,00         0,86 Hs.283681 ESTs           2031 GTGGCTCACGC         56,00         7,21 Hs.228230 EST           2032 CAGCTATTCA         10,00         1,69 Hs.153179 fatty acid binding protein 5 (psoria associated)           2033 CACACACACAC         4,00         0,84 Hs.63984 cadherin 13, H-cadherin (heart)           2034 CCACCACACAC         4,00         0,84 Hs.170222 solute carrier family 9 (sodium/hyd exchanger),           2035 CAAGGGTGACA         4,00         0,84 Hs.82035 GAP-like protein           2036 GCAGCACTTAT         2,00         0,48 Hs.7874 muskelin 1, intracellular mediator containing kelch           2039 AGCAGCCGCTC         2,00         0,48 Hs.76144 platelet-derived growth factor recebeta polype           2039 AGCAGCCGCTC         2,00         0,48 Hs.33039 ESTs           2041 TTATGCCTCCA         2,00         0,48 Hs.3304 ESTs	
Display	oinding
2027   AGCCACCACGC   17,00   2,63   Hs.60772   ESTs     2028   GCGAAACCTCG   8,00   1,46   Hs.210473   ESTs, Weakly similar to GELS_HI GELSOLIN PRECURSO     2029   GCAGTTGGATC   4,00   0,86   Hs.284932   Homo sapiens clone 24650 ubiqui hydrolase mRNA, p     2030   CTTGTGAAGTG   4,00   0,86   Hs.283681   ESTs     2031   GTGGCTCACGC   56,00   7,21   Hs.228230   EST     2032   CAGCTATTTCA   10,00   1,69   Hs.153179   fatty acid binding protein 5 (psoria associated)     2033   CACACACACAC   4,00   0,84   Hs.63984   cadherin 13, H-cadherin (heart)     2034   CCACCACACAC   4,00   0,84   Hs.256696   EST, Weakly similar to ALU1_HUI     ALU SUBFAMILY J SE     2035   CAAGGGTGACA   4,00   0,84   Hs.170222   solute carrier family 9 (sodium/hydexchanger),     2036   GCAGCACTTAT   2,00   0,48   Hs.82035   GAP-like protein     2037   CCTGACCTCAA   2,00   0,48   Hs.7874   muskelin 1, intracellular mediator     2039   AGCAGCGCTC   2,00   0,48   Hs.76144   platelet-derived growth factor received polype     2039   AGCAGCCGCTC   2,00   0,48   Hs.7104   Kruppel-like factor 13     2040   CTCCTGGCCCA   2,00   0,48   Hs.3314   ESTs     2041   TTATGCCTCCA   2,00   0,48   Hs.3314   ESTs     2042   TACTTCCTGCG   2,00   0,48   Hs.3314   Cas-Br-M (murine) ectropic retrovolation of transforming s     2044   GTAGGGTTCCT   2,00   0,48   Hs.278597   protein tyrosine phosphatase, non receptor type 18 (transposase-like protein     2045   ATAACCAAATG   2,00   0,48   Hs.278597   transposase-like protein	
2028   GCGAAACCTCG	
GELSOLIN PRECURSO	
hydrolase mRNA, p   2030 CTTGTGAAGTG	UMAN
2030 CTTGTGAAGTG         4,00         0,86 Hs.283681 ESTs           2031 GTGGCTCACGC         56,00         7,21 Hs.228230 EST           2032 CAGCTATTTCA         10,00         1,69 Hs.153179 fatty acid binding protein 5 (psoria associated)           2033 CACACACACAC         4,00         0,84 Hs.63984 cadherin 13, H-cadherin (heart)           2034 CCACCACACTC         4,00         0,84 Hs.256696 EST, Weakly similar to ALU1_HUI ALU SUBFAMILY J SE           2035 CAAGGGTGACA         4,00         0,84 Hs.170222 solute carrier family 9 (sodium/hyd exchanger),           2036 GCAGCACTTAT         2,00         0,48 Hs.82035 GAP-like protein           2037 CCTGACCTCAA         2,00         0,48 Hs.7874 muskelin 1, intracellular mediator containing kelch           2038 GGGGTATGGTT         2,00         0,48 Hs.76144 platelet-derived growth factor received polype           2039 AGCAGCCGCTC         2,00         0,48 Hs.7104 Kruppel-like factor 13           2040 CTCCTGGCCCA         2,00         0,48 Hs.3314 ESTs           2041 TTATGCCTCCA         2,00         0,48 Hs.38039 ESTs           2042 TACTTCCTGCG         2,00         0,48 Hs.3144 Cas-Br-M (murine) ectropic retrov transforming s           2044 GTAGGGTTCCT         2,00         0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (           2045 ATAACCAAATG         2,00         0,48 Hs.25726 transposon-derived Buster	itin
2031 GTGGCTCACGC         56,00         7,21 Hs.228230 EST           2032 CAGCTATTTCA         10,00         1,69 Hs.153179 fatty acid binding protein 5 (psorial associated)           2033 CACACACACAC         4,00         0,84 Hs.63984 cadherin 13, H-cadherin (heart)           2034 CCACCACACTC         4,00         0,84 Hs.256696 EST, Weakly similar to ALU1_HUI ALU SUBFAMILY J SE           2035 CAAGGGTGACA         4,00         0,84 Hs.170222 solute carrier family 9 (sodium/hydexchanger),           2036 GCAGCACTTAT         2,00         0,48 Hs.82035 GAP-like protein           2037 CCTGACCTCAA         2,00         0,48 Hs.7874 muskelin 1, intracellular mediator containing kelch           2038 GGGGTATGGTT         2,00         0,48 Hs.76144 platelet-derived growth factor receivet a polype           2039 AGCAGCCGCTC         2,00         0,48 Hs.7104 Kruppel-like factor 13           2040 CTCCTGGCCCA         2,00         0,48 Hs.5321 ARP3 (actin-related protein 3, year homolog           2041 TTATGCCTCCA         2,00         0,48 Hs.38039 ESTs           2042 TACTTCCTGCG         2,00         0,48 Hs.38039 ESTs           2043 CCTTGCCCAGG         2,00         0,48 Hs.38039 ESTs           2044 GTAGGGTTCCT         2,00         0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (           2045 ATAACCAAATG         2,00         0,48 Hs.25726 transposon-derived Buster1 t	
2032 CAGCTATTTCA 10,00 1,69 Hs.153179 fatty acid binding protein 5 (psorial associated)  2033 CACACACACAC 4,00 0,84 Hs.63984 cadherin 13, H-cadherin (heart)  2034 CCACCACACTC 4,00 0,84 Hs.256696 EST, Weakly similar to ALU1_HUI ALU SUBFAMILY J SE  2035 CAAGGGTGACA 4,00 0,84 Hs.170222 solute carrier family 9 (sodium/hydexchanger),  2036 GCAGCACTTAT 2,00 0,48 Hs.82035 GAP-like protein  2037 CCTGACCTCAA 2,00 0,48 Hs.7874 muskelin 1, intracellular mediator containing kelch  2038 GGGGTATGGTT 2,00 0,48 Hs.76144 platelet-derived growth factor received polype  2039 AGCAGCCGCTC 2,00 0,48 Hs.7104 Kruppel-like factor 13  2040 CTCCTGGCCCA 2,00 0,48 Hs.5321 ARP3 (actin-related protein 3, year homolog  2041 TTATGCCTCCA 2,00 0,48 Hs.43314 ESTs  2042 TACTTCCTGCG 2,00 0,48 Hs.38039 ESTs  2043 CCTTGCCCAGG 2,00 0,48 Hs.3144 Cas-Br-M (murine) ectropic retroved transforming s  2044 GTAGGGTTCCT 2,00 0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (  2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	
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2036 GCAGCACTTAT2,000,48 Hs.82035GAP-like protein2037 CCTGACCTCAA2,000,48 Hs.7874muskelin 1, intracellular mediator containing kelch2038 GGGTATGGTT2,000,48 Hs.76144platelet-derived growth factor receibeta polype2039 AGCAGCCGCTC2,000,48 Hs.7104Kruppel-like factor 132040 CTCCTGGCCCA2,000,48 Hs.5321ARP3 (actin-related protein 3, year homolog2041 TTATGCCTCCA2,000,48 Hs.43314ESTs2042 TACTTCCTGCG2,000,48 Hs.38039ESTs2043 CCTTGCCCAGG2,000,48 Hs.3144Cas-Br-M (murine) ectropic retrov transforming s2044 GTAGGGTTCCT2,000,48 Hs.278597protein tyrosine phosphatase, non receptor type 18 (2045 ATAACCAAATG2,000,48 Hs.25726transposon-derived Buster1 transposase-like protein	drogen
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containing kelch  2038 GGGGTATGGTT 2,00 0,48 Hs.76144 platelet-derived growth factor received polype  2039 AGCAGCCGCTC 2,00 0,48 Hs.7104 Kruppel-like factor 13  2040 CTCCTGGCCCA 2,00 0,48 Hs.5321 ARP3 (actin-related protein 3, year homolog  2041 TTATGCCTCCA 2,00 0,48 Hs.43314 ESTs  2042 TACTTCCTGCG 2,00 0,48 Hs.38039 ESTs  2043 CCTTGCCCAGG 2,00 0,48 Hs.3144 Cas-Br-M (murine) ectropic retrovitransforming s  2044 GTAGGGTTCCT 2,00 0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (  2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	
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2039 AGCAGCCGCTC2,000,48 Hs.7104Kruppel-like factor 132040 CTCCTGGCCCA2,000,48 Hs.5321ARP3 (actin-related protein 3, year homolog2041 TTATGCCTCCA2,000,48 Hs.43314ESTs2042 TACTTCCTGCG2,000,48 Hs.38039ESTs2043 CCTTGCCCAGG2,000,48 Hs.3144Cas-Br-M (murine) ectropic retrov transforming s2044 GTAGGGTTCCT2,000,48 Hs.278597protein tyrosine phosphatase, non receptor type 18 (2045 ATAACCAAATG2,000,48 Hs.25726transposon-derived Buster1 transposase-like protein	eptor,
2040 CTCCTGGCCCA 2,00 0,48 Hs.5321 ARP3 (actin-related protein 3, year homolog  2041 TTATGCCTCCA 2,00 0,48 Hs.43314 ESTs  2042 TACTTCCTGCG 2,00 0,48 Hs.38039 ESTs  2043 CCTTGCCCAGG 2,00 0,48 Hs.3144 Cas-Br-M (murine) ectropic retrov transforming s  2044 GTAGGGTTCCT 2,00 0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (  2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	
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2042 TACTTCCTGCG2,000,48 Hs.38039ESTs2043 CCTTGCCCAGG2,000,48 Hs.3144Cas-Br-M (murine) ectropic retrov transforming s2044 GTAGGGTTCCT2,000,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (2045 ATAACCAAATG2,000,48 Hs.25726 transposon-derived Buster1 transposase-like protein	ast)
2043 CCTTGCCCAGG 2,00 0,48 Hs.3144 Cas-Br-M (murine) ectropic retrover transforming s  2044 GTAGGGTTCCT 2,00 0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (  2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	
transforming s  2044 GTAGGGTTCCT 2,00 0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 (  2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	
2044 GTAGGGTTCCT 2,00 0,48 Hs.278597 protein tyrosine phosphatase, non receptor type 18 ( 2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	riral
receptor type 18 ( 2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	า-
2045 ATAACCAAATG 2,00 0,48 Hs.25726 transposon-derived Buster1 transposase-like protein	
transposase-like protein	
2046 CCTGTGAATAG 2,00 0,48 Hs.200647 EST	
2047 TGACCAGGGTC 2,00 0,48 Hs.167827 Homo sapiens clone HH419 unkn	iown
2048 CAGACCCAAAA 2,00 0,48 Hs.167558 zinc finger protein 161	

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2050	TTCTCATAATC	2,00	0,48	IHs.164919	ESTs, Highly similar to KPC2_HUMAN
					PROTEIN KINASE C,
2054	<u>ACTGATGCAAG</u>	2,00	0,48	Hs.161049	ESTs
	AACTCTGATAT	2,00	0,48	Hs.151046	hypothetical protein FLJ11193
2052	GGGCATCTCCA	2,00	0,48	Hs.107000	ESTs
2053	TGGATGTCTGT	1,00	0,24	Hs.96716	ESTs
2054	TATTAGAAGCA	1,00	0,24	Hs.91065	hypothetical protein DKFZp761B2423
2055	CACTGAATATG	1,00	0,24		small nuclear ribonucleoprotein D1 polypeptide (16kD
2056	AGTTAATAAAG	1,00	0,24	Hs.8065	Homo sapiens mRNA full length insert cDNA clone EURO
2057	TTATATTTTCT	1,00	0,24	Hs.8021	KIAA1058 protein
2058	GTCACCAAACA	1,00			selectin P ligand
	TTACACTGTAA	1,00	0,24		neutral sphingomyelinase (N-SMase) activation associ
	AAGTTGCATCT	1,00	0,24	Hs.74649	cytochrome c oxidase subunit VIc
_	TAGACATTTGA	1,00			cytochrome c oxidase subunit VIc
2062	TTCATAGGTAA	1,00	0,24	Hs.7312	ESTs
2063	CATTTTAGGCA	1,00	0,24	Hs.72782	hypothetical protein FLJ11171
2064	ATACTATAATT	1,00	0,24		Human DNA sequence from clone RP1- 187J11 on chromoso
2065	CTTCGCTTTGT	1,00	0,24	Hs.69485	ESTs, Weakly similar to similar to other protein pho
2066	ATGATATATGC	1,00	0,24	Hs.6831	Homo sapiens clone 1400 unknown protein mRNA, partia
2067	GCCCCCTTACA	1,00	0,24	Hs.49476	Homo sapiens clone TUA8 Cri-du-chat region mRNA
2068	TCAGTCCCTGT	1,00	0.24		hypothetical protein FLJ20209
2069	AATTITCATTA	1,00			ESTs
2070	GCCAGTCCACT	1,00		Hs.34782	ESTs, Moderately similar to transducin [H.sapiens]
2071	TAGAGAGTTTA	1,00	0,24	Hs.29643	
2072	GCTGTCCCCTC	1,00	0,24	Hs.278422	DKFZP586G1122 protein
2073	GTGAAAGCCGT	1,00		Hs.270662	ESTs, Weakly similar to transformation- related prote
	TCTTCCAGAAA	1,00	0,24	Hs.256585	
2075	TTGGCCGGGAT	1,00		Hs.254900	
	GTAAAGAATGT	1,00			ESTs
2077	AAATTTTTGTA	1,00		Hs.24650	ESTs, Moderately similar to AF133913_1 ARL-6 interac
2078	AACGCTGCAAA	1,00	0.24		KIAA0876 protein
	GTACCCTAAAA	1,00		Hs.239970	ESTs, Weakly similar to b34l8.1 [H.sapiens]
2080	TAATCTTTTTT	1,00	0.24	Hs.231463	
	AAATTGTATGT	1,00			tropomodulin 3 (ubiquitous)
	CCACTACATTC	1,00			ESTs
	GCCGCACTCAG	1,00		Hs.200577	
	TCTTGTCATAC	1,00			conserved helix-loop-helix ubiquitous

	CACCACCTCAT	4.00	004	11- 400004	
	CAGCACCTGAT	1,00			pyruvate kinase, muscle
	TTGAATAAAAG	1,00			phospholipase A2, group IVB (cytosolic)
2087	GGGAAGTGTGC	1,00	0,24	HS.197733	ESTs, Weakly similar to TRP7_HUMAN TRANSIENT RECEPTO
2088	CCTGGCCTACC	1,00	0,24	Hs.19585	KRAB-zinc finger protein SZF1-1
2089	TITATTTTCAA	1,00	0,24	Hs.194293	ESTs
2090	TCTCTCTGCCT	1,00	0,24	Hs.184987	ESTs
2091	AGAGGAAGTAA	1,00	0,24	Hs.177537	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
2092	ACCAAAAAAA	1,00	0,24	Hs.173724	creatine kinase, brain
2093	GGGACATTTAT	1,00	0,24	Hs.173108	Homo sapiens clone 24523 mRNA sequence
2094	TACCATCCATA	1,00	0,24	Hs.169476	glyceraldehyde-3-phosphate dehydrogenase
2095	TTGATGAAGAA	1,00	0,24	Hs.168075	karyopherin (importin) beta 2
	GCCCGGTGCCC	1,00		Hs.1665	zinc finger protein homologous to Zfp-36 in mouse
2097	AAGAACTGTTT	1,00	0 24	Hs 159456	zinc finger protein 288
_	CCAGCAACTGT	1,00			SET translocation (myeloid leukemia-
					associated)
	ATGGTGCGTGC	1,00			ESTs, Weakly similar to ALUE_HUMAN !!!! ALU CLASS E
2100	CTCTTCAGGGT	1,00	0,24	Hs.13781	Homo sapiens cDNA FLJ11302 fis, clone PLACE1009971
2101	TCAATGTGAAA	1,00	0,24		Homo sapiens BAC clone RP11-121A8 from 7p14-p13
2102	CTTAATACTAC	1,00	0,24		KIAA0592 protein
2103	GGCTGCAGTAT	1,00			KIAA0522 protein
2104	GGAAGCTGAAG	1,00		Hs.128629	
2105	CCTCGGGCATC	1,00		Hs.126735	
2106	тттсттвств	1,00			Homo sapiens mRNA; cDNA DKFZp547D135 (from clone DKF
2107	GCCTTGCCTCT	1,00	0,24	Hs.118837	ESTs
2108	CTTTAAAAAAA	1,00			fibronectin 1
2109	TGCGGAAAAA	1,00			G protein-coupled receptor 30
	TGACTTTCTGC	1,00	0,24	Hs.11123	ESTs, Weakly similar to B38919 hypothetical protein
2111	CATTTACATAT	1,00	0,24		Homo sapiens clone 24775 mRNA sequence
2112	AATGTGTTACT	1,00	0.24	Hs.105751	Ste20-related serine/threonine kinase
	ATCTTTATTCC	1,00			KIAA0308 protein
	TGTTTGAATTC	1,00			Homo sapiens mRNA; cDNA DKFZp434F1622 (from clone DK
2115	ACAACACCCCA	5,00	1,00	Hs.21453	Homo sapiens mRNA for inositol 1,4,5- trisphosphate 3
2116	CTGGAAATAAA	3,00	0.67	Hs.69745	ferredoxin reductase
	TTATTTATGAA	3,00			tissue inhibitor of metalloproteinase 3 (Sorsby fund
2118	TACCAAGGATT	3,00	0.67	Hs.21729	splicing factor 3a, subunit 1, 120kD
<u> </u>	MOURAGOATT	3,00	0,07	113.21/23	phioning ractor sa, subullit 1, 120kD

2110	GTGCCAAACAC	2.00	0.67	Un 172216	Johannesenia A (norothyroid popular)
2119	GIGCCAAACAC	3,00	0,67	MS. 1/2216	chromogranin A (parathyroid secretory protein 1)
2120	CCCGGCCCAAA	3,00	0.67	Hs.133207	PTPRF interacting protein, binding
					protein 1 (liprin
2121	AGAATTGCTTG	37,00	4,88	Hs.56542	X-prolyl aminopeptidase
L					(aminopeptidase P) 1, solubl
	TCTCTGATGCT	34,00		Hs.6441	tissue inhibitor of metalloproteinase 2
2123	ACACTGCACTC	7,00	1,27	Hs.200454	ESTs, Weakly similar to ALU1_HUMAN
0404	04447040040			=====	ALU SUBFAMILY J S
_	GAAATGAGCAG	6,00			KIAA0127 gene product
	GCTGGATGCGG	6,00		Hs.18075	chromosome 9 open reading frame 3
	GCAAAACCCTA	6,00			DKFZP586A0522 protein
	TTTGCTCTCCC	15,00		Hs.75350	vinculin
	GGAGGCTGAGG	48,00			membrane fatty acid (lipid) desaturase
	GTAAAACCCCA	21,00		Hs.18955	Homo sapiens cDNA FLJ20667 fis, clone KAIA596
	GGAGGGGCTT	30,00			lamin A/C
2131	AGCTAAGTTTG	4,00	0,83	Hs.19447	ESTs, Weakly similar to CL36_HUMAN LIM DOMAIN PROTEI
2132	CACACAGTTTT	18,00	2,62	Hs.204354	ras homolog gene family, member B
2133	ATGGCGGGTGC	9,00			hypothetical protein FLJ20001
2134	GACTTGTATAT	6,00			nuclear factor of kappa light polypeptide
					gene enhan
2135	ATAGTACAGCC	3,00			MEK partner 1
	AGCCACCGCTC	_3,00		Hs.6195	ESTs
2137	GGCAGGATGAT	3,00	0,66	Hs.274319	hypothetical protein FLJ10509
	GCCTGGGACCT	3,00	0,66	Hs.180871	protein kinase C, alpha binding protein
2139	ACAGCCGTGGG	3,00	0,66	Hs.123090	SWI/SNF related, matrix associated, actin dependent
2140	AGCCGAGATCA	5,00	0,97	Hs.277663	EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SE
2141	AGCCTTTGTTG	26,00	3,47	Hs.9930	collagen-binding protein 2 (colligen 2)
2142	GTGTGCCTCCA	6,00			interferon regulatory factor 3
2143	TGGAGAAGAGC	8,00	1,37		upregulated by 1,25-dihydroxyvitamin D-
2144	GGCCCCATTGC	4,00	0,81		Human clone CE29 8.1 (CAC)n/(GTG)n repeat-containing
2145	AATATTCATAG	2,00	0,47	Hs.8583	similar to APOBEC1
	GCTCCGTAAGG	2,00		Hs.80712	KIAA0202 protein
	GAGGTGCCCCA	2,00			ESTs
	AGTGTATTTT	2,00			insulin-like growth factor 2 receptor
	TAAACCTAAAG	2,00		Hs.60548	hypothetical protein PRO1635
2150	AGTCAAGCCCC	2,00			four and a half LIM domains 3
2151	TATCAAAACAT	2,00		Hs.258939	
	ATCCTACTGTT	2,00			uncharacterized hypothalamus protein HCDASE
2153	GCACCAAATGA	2,00	0.47	Hs.23585	KIAA1078 protein
	CCAATGCTATG	2,00		Hs.22753	Homo sapiens mRNA; cDNA
		_,,,,			DKFZp434K0926 (from clone DK

2155	CATCGTTACAT	2,00	0.47	Hs.173802	KIAA0603 gene product
	TAACCAAAAAC	2,00			ELK4, ETS-domain protein (SRF
		_,	-,		accessory protein 1)
2157	CTGAAACCCCA	2,00	0,47	Hs.162132	
	GGAAGGCAAGC	2,00			ESTs, Weakly similar to ALU7_HUMAN
		· 1	•		ALU SUBFAMILY SQ
2159	GAAAGGTGGTT	2,00	0,47	Hs.14394	hypothetical protein FLJ20157
2160	TCATAACCTTG	2,00	0,47	Hs.124029	inositol polyphosphate-5-phosphatase,
					40kD
	TACCCCATAAA	5,00		Hs.281083	
2162	CGCCCCCTGCG	6,00	1,09	Hs.135805	ESTs, Weakly similar to KIAA1323 protein [H.sapiens]
2163	ACAAAGCCCCA	3,00	0,65	Hs.8583	similar to APOBEC1
2164	CCTATAGTCCT	3,00	0,65	Hs.41694	origin recognition complex, subunit 2
					(yeast homolog
2165	CCAGTACAGCC	3,00	0,65	Hs.140978	Homo sapiens mRNA; cDNA
					DKFZp762H106 (from clone DKF
	CTGCTGCTGGT	3,00			Cdc42 effector protein 2
	CTTAATCTTGT	4,00			BTG family, member 2
2168	GCCACACCCCC	4,00	0,80	Hs.113916	Burkitt lymphoma receptor 1, GTP-
2422					binding protein
	TCCTCCCTACT	14,00			yeast Sec31p homolog
	GTGAGACCCCA	12,00		Hs.198671	
	CAGATGCAAAA	11,00			paired box gene 6 (aniridia, keratitis)
21/2	CCCCAGGAGAA	5,00	0,93		solute carrier family 2 (facilitated glucose
2472	CTTOTOCOCOAO	0.00	4.00		transpo
21/3	CTTCTGGGGAC	6,00	1,06	Hs.75082	ras homolog gene family, member G (rho G)
2174	GTGGCTTACAC	4,00	0.79	He 264482	Homo sapiens mRNA; cDNA
2117	01000117070	7,00	0,73	113.20-1-102	DKFZp761A0411 (from clone DK
2175	GCGAGACCCCA	4,00	0.79	Hs.15681	ESTs
	TAAACTATTGG	3,00			KIAA0217 protein
$\overline{}$	GACGGCTGCAA	3,00		Hs.4909	dickkopf (Xenopus laevis) homolog 3
	GTGAAACCGTC	3,00			Homo sapiens mRNA full length insert
		,,,,,			cDNA clone EURO
2179	CCACTGGACTC	3,00	0,63		ESTs, Weakly similar to ALU1_HUMAN
					ALU SUBFAMILY J S
	TGTTAGCAAAT	3,00		Hs.22666	
	AAGTACGAGGA	3,00		Hs.22660	ESTs
	CTGTTGGAAAA	3,00		Hs.209863	
2183	AACCACTGTGC	3,00	0,63	Hs.188037	ESTs, Moderately similar to
					ALU7_HUMAN ALU SUBFAMILY
	CTTCTCCAAAA	2,00		Hs.99949	prolactin-induced protein
	GTGGTCAAGTT	2,00		Hs.92127	ESTs
2186	CCTTTGCTGAG	2,00	0,46	Hs.7442	Human DNA sequence from clone
24.5=					742C19 on chromosome 2
2187	TTGCTGCCAGC	2,00	0,46	Hs.5566	gap junction protein, beta 2, 26kD
0400	TOAATTTOOTO	0.00	6.45	11 00055	(connexin 26)
2188	TGAATTTCCTG	2,00	<u>0,46</u>	Hs.30057	Homo sapiens clone 24749 and 24750

			_		mPNA sogueness
2490	CTCCTATATCC	2.00	0.46		mRNA sequences
	GTGGTATATGC	2,00			hypothetical protein FLJ20342
2190	GTGGTGCACTC	2,00	0,46	HS.2000/3	Homo sapiens mRNA; cDNA
2101	CTCCCACAACC	2.00	0.46	U- 250407	DKFZp762N226 (from clone DKF
$\overline{}$	GTGGCACAAGC	2,00		Hs.258487	
	AGCCACCATAC	2,00		Hs.236051	
2193	TAAAACCGTTT	2,00	0,46	Hs.182280	MADS box transcription enhancer factor
0404	00770777440		0.40	11 470007	2, polypeptid
2194	CCTTGTTTAAC	2,00	0,46	Hs.1/3965	ribosomal protein S6 kinase, 90kD,
2405	COTOAOTOOAA	- 0 00	0.40	U- 400005	polypeptide 3
	GCTCACTGCAA	2,00		Hs.163385	
	TGCCTTGGGCT	2,00			retinoic acid receptor, gamma
	TTATTGTTCCC	2,00			hepatocellular carcinoma-associated antigen 112
	GGGACAACCCA	2,00			ESTs
-	GTTTCTTACTG	2,00			KIAA1327 protein
	TTTTGCTTTTT	2,00			lysyl oxidase
2201	ATTITCATCAA	2,00	0,46	Hs.101282	Homo sapiens mRNA; cDNA
					DKFZp434B102 (from clone DKF
2202	TACAGAATGTG	1,00			ESTs
2203	TTGTATTGTTG	1,00			ESTs
	GGGCAGAAGGC	1,00	0,23	Hs.9585	ESTs
2205	TATTGGCCTGG	1,00	0,23	Hs.79572	cathepsin D (lysosomal aspartyl protease)
2206	TGACTGTAAAA	1,00	0,23	Hs.75621	protease inhibitor 1 (anti-elastase), alpha-1-antitr
2207	ACCTGCTTCCC	1,00	0.23	Hs.75458	ribosomal protein L18
	TGATACAGAAA	1,00			ESTs
	AATGAATTCTT	1,00			Homo sapiens mRNA; cDNA
1 1					DKFZp564E2222 (from clone DK
2210	ATTITGTCACT	1,00	0,23	Hs.5459	KIAA1436 protein
	CACCCTTTACT	1,00		Hs.5437	Tax1 (human T-cell leukemia virus type I) binding pr
2212	CTGGTGGGCCA	1,00	0.23	Hs.5338	carbonic anhydrase XII
	CTTTGAAATAG	1,00			ESTs
2214	AGTCCTGCTTC	1,00			ESTs
	TAACATTGAGA	1,00			ESTs
	CTGAGAAGCGG	1,00		Hs.41055	ESTs, Highly similar to RL2A_HUMAN 60S RIBOSOMAL PRO
2217	TATATATAGAG	1,00	0,23	Hs.34853	inhibitor of DNA binding 4, dominant
					negative helix-
2218	GTCTGCTCCAG	1,00	0,23	Hs.32978	proprotein convertase subtilisin/kexin type 7
2219	CCCCGAGGCT	1,00	0,23	Hs.31019	ESTs
	AATGAAAAATT	1,00			cytochrome c oxidase subunit VIIa polypeptide 2 like
2221	TTTGTACTAAT	1,00	0,23	Hs.29846	Human DNA sequence from clone 717M23 on chromosome 2
2222	TTCCATTATCA	1,00	0.23	Hs.29356	ESTs
		.,00	<u> </u>		<u> </u>

					r
	GTTTAAAAAGC	1,00		Hs.29032	
	CAATITAAAGT	1,00		Hs.286249	
	TGTATTTGTAA	1,00	0,23	Hs.286056	ESTs
2226	GTGGCGTGCCT	1,00		Hs.282652	
2227	TAAAATGTGAC	1,00	0,23	Hs.268447	ESTs, Weakly similar to ALU8_HUMAN ALU SUBFAMILY SX
2228	AACAATAAAAA	1,00	0,23		ESTs, Weakly similar to ALU4_HUMAN ALU SUBFAMILY SB2
2229	GAAACACGTAG	1,00	0,23	Hs.26407	ESTs
2230	ACTITAACAGG	1,00	0,23		cadherin EGF LAG seven-pass G-type receptor 1
2231	AGCCCCACAAA	1,00	0,23	Hs.250570	ESTs
2232	TGGTGAGATGA	1,00	0,23	Hs.224829	ESTs
	GGCACCGCGTG	1,00	0,23	Hs.20677	KIAA1303 protein
$\overline{}$	CAAGGATAAGA	1,00		Hs.2001	thromboxane A synthase 1 (platelet, cytochrome P450,
2235	CGCCCGGCGG	1,00	0,23	Hs.196244	ESTs
2236	GAAGAGAAGGT	1,00	0,23	Hs.180455	RAD23 (S. cerevisiae) homolog A
2237	CAGATTTTGTG	1,00	0,23		calmodulin 1 (phosphorylase kinase, delta)
2238	ATTAGTTACAA	1,00	0,23	Hs.177635	KIAA1095 protein
2239	CCCACCACATT	1,00		Hs.170610	
	CTGCTAACCCA	1,00			cat eye syndrome chromosome region, candidate 1
2241	GTTCTCTGCTT	1,00	0,23		Human clone A9A2BRB6 (CAC)n/(GTG)n repeat-containing
2242	TATGTACAGTT	1,00	0,23		hypothetical protein PRO0907
2243	AACCCGGGAGT	1,00		Hs.161974	
2244	GTAACCTCAAA	1,00	0,23	Hs.161930	EST
2245	TTCAATTTCTT	1,00	0,23		SWI/SNF related, matrix associated, actin dependent
2246	AGCTAAAAAAA	1,00	0,23	Hs.15953	hypothetical protein FLJ10120
2247	TTATTTAAAAG	1,00			KIAA0981 protein
2248	GACTCCACATT	1,00		Hs.156637	Cas-Br-M (murine) ectropic retroviral transforming s
2249	AGACAAAATTA	1,00	0,23	Hs.153487	signal transducing adaptor molecule (SH3 domain and
2250	ACTGAGAAGAG	1,00	0,23	Hs.130761	ESTs, Moderately similar to AF151803_1 CGI-45 protei
2251	GGTGCTGAATA	1,00	0,23		ESTs
	GATTTCTTTGA	1,00		Hs.119651	
	TGTCAAAAGAG	1,00			Homo sapiens cDNA FLJ20053 fis, clone COL00809
2254	ACGGGGAGAGT	1,00	0.23	Hs.117582	CGI-43 protein
	TGGCTTTATCC	1,00			Human clone 23589 mRNA sequence
	GTTGCGTGTCC	1,00			NOT3 (negative regulator of transcription 3, yeast)
2257	TAAACGGCCTC	_1,00	0.23	Hs.10632	hypothetical protein DKFZp762M136
	GGCCAGAATGA	1,00			DKFZP434A043 protein
	- JOURNATUR			1.13.132130	DIVIET TOTI (010 PIOCOIII

2250	TGAAGCAAAAA	1.00	0.22	Ha 100407	Hama caniona mRNA: aDNA
2259	IGAAGCAAAAA	1,00	0,23	ms. 100407	Homo sapiens mRNA; cDNA DKFZp564H2416 (from clone DK
2260	TCACAGCTGTG	19,00	2.52	Hs.77054	B-cell translocation gene 1, anti-
2200	TOACAGCTGTG	19,00	2,52	113.77034	proliferative
2261	AGAAAAAAAA	70,00	7.56	Hs 251680	(Manual assignment) not unique,
		. 0,00	.,		contains loricrin
2262	GTGGCGCACGC	11,00	1,65	Hs.135723	glycolipid transfer protein
2263	TCTGTACACCT	19,00	2,51	Hs.182740	ribosomal protein S11
2264	AGGGAGGGCC	6,00	1,05	Hs.172153	glutathione peroxidase 3 (plasma)
2265	AGGACACCGCC	5,00	0,91	Hs.77793	c-src tyrosine kinase
2266	CGAGGGCACTC	5,00			spectrin, beta, non-erythrocytic 2
	TGCTGCCAGAC	4,00	0,77	Hs.79219	RalGDS-like gene; KIAA0959 protein
2268	TACTGGTTTAT	4,00	0,77	Hs.30299	IGF-II mRNA-binding protein 2
2269	CCCCCACCTAA	15,00	2,04	Hs.77422	proteolipid protein 2 (colonic epithelium-
					enriched)
	TGAATGATACG	4,00			protease, serine, 15
-	TGAAACTGCAA	4,00		Hs.147189	HYA22 protein
	TGAGTCTGGCT	10,00		Hs.4055	chromosome 21 open reading frame 50
2273	GGAGTGTGCTC	29,00	3,36	Hs.9615	myosin regulatory light chain 2, smooth muscle isofo
2274	TCATCGGGCTG	2,00	0,44	Hs.78335	microtubule-associated protein, RP/EB
2275	CACACOTTOCA	2.00	0:44	11- 70040	family, member
	GAGACCTTGGA	2,00		Hs.72249	protease-activated receptor 3
	CACTGGACGAG	2,00			ESTs
2211	TGTACTTATTA	2,00	0,44	Hs.6906	v-ral simian leukemia viral oncogene
2279	ACCACCCTGTT	2,00	0.44	Hs.4864	homolog A (ras KIAA0892 protein
	ATTICTCATTC	2,00			
	GGACATTTTC	2,00			D-type cyclin-interacting protein 1
	CATCTTAAATG	2,00			hypothetical protein FLJ11046 hypothetical protein FLJ20725
	GAATCATTTAT	2,00			KIAA0391 gene product
	ATCAAATGCAA	9,00		Hs.79070	v-myc avian myelocytomatosis viral
2200	ATOMATOMA	3,00	1,30	115.73070	oncogene homolog
2284	AGCACCAGAAC	3,00	0.61	Hs.60103	KIAA0690 protein
	TCCTGACCACC	3,00			LIM domain binding 1
	CTTATAATCCC	3,00		Hs.259541	
	ACCCATCGCCT	3,00		Hs.165428	
2288	CCATTGCACTA	3,00			ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
2289	TACCCCAGAAC	9,00	1.35	Hs.145320	
	CTCTCACCCTG	18,00		į	ribonuclease/angiogenin inhibitor
	CAAGGCTTGC	15,00			RAP1B, member of RAS oncogene
					family
2292	TTTGCACTTGT	7,00	1,12	Hs.75188	wee1+ (S. pombe) homolog
	GCCAGGAGCTA	7,00	1,12	Hs.18141	ladinin 1
2294	TAAAATACTCC	4,00	0,75	Hs.8125	Homo sapiens mRNA; cDNA
220F	TGATTGATTTC	4.00	0.75	Un 5040	DKFZp586E1521 (from clone DK
	TOANAGETTE	4,00		Hs.5912	F-box only protein 7
2280	TAAAAACTTTC	4,00	0,75	ms.204096	lipophilin B (uteroglobin family member),

$\overline{}$	<del> </del>	TT		ī	Innestatoin
2207	TOACTOCACTO	16.00	2 00	U- 004000	prostatein
	TCACTGCACTC	16,00		Hs.261038	
	ACCAAAGCCCC	10,00			Human putative ribosomal protein S1 mRNA
2299	AATAGGGTCAA	8,00	_		amyloid beta (A4) precursor-like protein 2
2300	CCTATAGTCCC	11,00	1,53	Hs.140697	ESTs, Weakly similar to unnamed protein product [H.s
2301	CCTCCCTGCTC	3,00	0,59	Hs.90790	ESTs
2302	ACCTAGCCACT	3,00	0,59	Hs.89463	potassium large conductance calcium- activated channe
2303	CAGCTCAGCTG	3,00			filamin C, gamma (actin-binding protein- 280)
2304	TCAATAAATGT	3,00	0,59	Hs.106747	ESTs, Weakly similar to AF217508_1 uncharacterized b
2305	CACTCACACCC	4,00	0,73	Hs.24447	sigma receptor (SR31747 binding protein
2306	GTGAAACCTGT	7,00	1,10	Hs.272795	hypothetical protein FLJ20359
	ATCATACCACG	2,00	2,80	Hs.97259	ESTs
	AACGGGGCCCT	2,00			small inducible cytokine subfamily A
		'			(Cys-Cys), memb
2309	TCCCCGGTCAG	2,00	2,80	Hs.80562	gelsolin (amyloidosis, Finnish type)
2310	GCGCATCAAAA	2,00			ESTs
2311	AATTTCAGGCA	2,00	2,80	Hs.5476	serine protease inhibitor, Kazal type, 5
2312	ATCCCACTACT	2,00			ESTs
2313	ATCCGCTGGGG	2,00	2,80	Hs.30954	phosphomevalonate kinase
2314	ACTGTGGACTG	2,00	2,80	Hs.285122	ESTs, Weakly similar to S53869 laminin beta-2 chain
2315	AGAGAAGAATG	2,00	2,80	Hs.2841	neuromedin U
2316	AGGATTGTTTG	2,00	2,80	Hs.283545	
2317	GTAAAGATTTG	2,00		Hs.278629	
2318	ACTATGGATAG	2,00		Hs.275511	
2319	GTTAGGCACGA	2,00	2,80	Hs.272800	hypothetical protein FLJ20456
	TGCCACCAACA	2,00			beta-site APP-cleaving enzyme 2
	AAATGGGAACA	2,00		Hs.271226	
	GGGGTTTGTTT	2,00		Hs.258455	
2323	ACATAGTCTGA	2,00		Hs.25766	
	TATGCTGAAAT	2,00		Hs.255277	
	TTCACTCCATT	2,00		Hs.254914	EST, Weakly similar to NICE-1 protein [H.sapiens]
2326	TCTTGGCAACA	2,00	2.80		KIAA1219 protein
_	CAATAAAATTC	2,00		Hs.250236	
	GGCTTTGGTT	2,00		Hs.230730	EST, Moderately similar to RLA1_HUMAN 60S ACIDIC RIB
2329	AGTTTGTTTTA	2,00	2.80	Hs.212570	
	TTCCACTGTGA	2,00		Hs.198862	
	CAAGTGGGTGT	2,00		Hs.187685	
	TGCAGGGACCT	2,00			metastasis-associated 1-like 1
	TACTTCACCCA	2,00			aldehyde dehydrogenase 5

2334	CTTTGATTTAT	2,00	2.80	Hs.165590	ribosomal protein S13
	TGTTGTTTTT	2,00			Homo sapiens mRNA; cDNA
		_,,55	_,00		DKFZp434K1111 (from clone DK
2336	GTAGCGCCTCC	2,00	2.80	Hs 143212	cystatin F (leukocystatin)
	CTAGTGGCGGC	2,00			ESTs, Weakly similar to ALU8_HUMAN
					ALU SUBFAMILY SX
2338	ACCTCACCTGG	2,00	2,80	Hs.137585	UDP glycosyltransferase 2 family, polypeptide B11
2339	CTTGTTCAAAA	2,00	2.80	Hs 137560	ESTs, Moderately similar to AMSH
					[H.sapiens]
	CCTCTTTCCAG	2,00		Hs.134615	
	TCTTAGTTCTA	2,00		Hs.130729	
	CACCTTATAGT	2,00			CGI-43 protein
2343	AGCTTCTACCA	2,00			small proline-rich protein 2A
2344	GTCAAGCCCAA	2,00	2,80		ESTs, Weakly similar to SPR2J protein [M.musculus]
2345	AACTTATCATT	2,00	2,80	Hs.103368	
	ATGTGCTTCCG	2,00			proline arginine-rich end leucine-rich repeat protei
2347	CCTTTTGTCC	2,00	0.43	Hs.62601	Homo sapiens mRNA; cDNA
-0		_,00	0,40	110.02001	DKFZp586K1318 (from clone DK
2348	CTCTGTTTACA	2,00	0,43	Hs.5947	mel transforming oncogene (derived from cell line NK
2349	GAAGTGCTGCT	2,00	0.43	Hs.21812	
	CCATTGCAGTC	2,00		Hs.207659	
	CCACTGCACGC	2,00			thiopurine S-methyltransferase
	GTGGTGTACAC	2,00			Human proteinase activated receptor-2
					mRNA, 3'UTR
	ATAAATAAATT	2,00			hypothetical protein FLJ10506
	TTGATAAATAA	2,00			replication factor C (activator 1) 2 (40kD)
	GCAAAAGCCCG	2,00		Hs.109798	
	TAAAATATGGG	1,00	0,22	Hs.98401	Homo sapiens mRNA full length insert cDNA clone EURO
2357	CTTTGGTTTGC	1,00	0,22	Hs.8895	ESTs
2358	CCAAAGGAGAA	1,00	0,22	Hs.8889	serine hydroxymethyltransferase 1 (soluble)
2359	TGTACCTTTTC	1,00	0,22	Hs.8172	ESTs
	TGTTAAGTGTT	1,00		Hs.78825	matrin 3
	TACAAGCTGAG	1,00		Hs.77508	glutamate dehydrogenase 1
	TCTGCGGGTGG	1,00		Hs.76159	ATPase, H+ transporting, lysosomal
					(vacuolar proton
2363	TATCAATATTC	1,00	0,22	Hs.7306	secreted frizzled-related protein 1
2364	ATGTTTTGTAA	1,00		Hs.6853	carbohydrate (N-acetylglucosamine 6-0) sulfotransfer
2365	GCTTTTTGTTG	1,00	0,22	Hs.6048	FEM-1-like death receptor binding protein
2366	CTGGAGACCCA	1,00	0 22	Hs.56729	lymphocyte-specific protein 1
	TTCAGAATAAA	1,00			hypothetical protein FLJ20331
					<del></del>
2308	ACAATGAAGCA	1,00	0,22	HS.285665	Homo sapiens mRNA; cDNA

2369  GCCTGTGGGGT						
2370   CAGCAAGGCTT						DKFZp564P013 (from clone DKF
2371   CAAGGCACCAA						
2372   TACCAGAGTCC   1,00   0,22   Hs.279927   hypothetical protein containing pH domain, PTB domain and PTB						
2373   ATTITIGTAAA						
PTB domain and	2372	TACCAGAGTCC	1,00		Hs.279927	hypothetical protein
2374   TCTGTTACACC	2373	ATTTTGTAAA	1,00	0,22	Hs.27413	
2375   AGGTTTGATTT	2374	TCTGTTACACC	1,00	0,22	Hs.272759	
2376   AAGAGATGTTC						
2377   TTGGTCAGGGT			_			
1,00						
2379   AGACTCAGGCC   1,00   0,22   Hs.24305   ESTs						
2380 GTTCATCCTTG	_					
2382   GTTTGTTTCCT					Hs.236894	ESTs, Highly similar to LRP1_HUMAN
protein [H.sapiens]   protein [H.sapiens]   2388 AAGACACTGTT   1,00   0,22   Hs.20707   hypothetical protein R31240_1   2385 GTCGGGGGAGA   1,00   0,22   Hs.18844   ESTs   2386 ATGTTGTCAAT   1,00   0,22   Hs.18844   ESTs   MHC class I region ORF   2387 TTTTCCCTCAG   1,00   0,22   Hs.184242   sterol-C5-desaturase (fungal ERG3, delta-5-desaturas   2388 ACATTCATTA   1,00   0,22   Hs.18192   Ser/Arg-related nuclear matrix protein (plenty of pr   2389   TTAGCCAGGGT   1,00   0,22   Hs.180610   splicing factor proline/glutamine rich (polypyrimidi   2390   GGTGGAAAAAA   1,00   0,22   Hs.1878728   methyl-CpG binding domain protein 3   2391   GGGCAGAATAA   1,00   0,22   Hs.164690   ESTs   2392   GGTGCCCGGCA   1,00   0,22   Hs.164690   ESTs   2393   TAGCTGAGGCA   1,00   0,22   Hs.164690   ESTs   2394   CGGATTCAATT   1,00   0,22   Hs.164690   ESTs   2395   AAGAAAGGGAA   1,00   0,22   Hs.14763   ESTs   2395   AAGAAAGGGAA   1,00   0,22   Hs.14908   ESTs   2396   TCAAAAGGGCA   1,00   0,22   Hs.14908   ESTs   2397   CTCTCCTTGCC   1,00   0,22   Hs.16913   ESTs   2398   TGCAGAGAGG   1,00   0,22   Hs.10373   ESTs   2399   TGTCGCAGAA   1,00   0,22   Hs.107418   ESTs   2400   AGCTTCCGGT   1,00   0,22   Hs.107418   ESTs   2400   AGCTTCCGAG   15,00   1,89   Hs.23582   tumor-associated calcium signal transducer 2   2403   GCCCACAGTAG   4,00   0,72   Hs.76122   splicing factor, arginine/serine-rich 4   2404   ACTGCAGAGCG   4,00   0,72   Hs.76122   splicing factor, arginine/serine-rich 4   2404   ACTGCAGAGCG   4,00   0,72   Hs.76122   splicing factor, arginine/serine-rich 4   2405   GGCCCCTCACC   11,00   1,48   Hs.274313   insulin-like growth factor binding protein   6	2381	TATCTTGTTGC	1,00	0,22		
2384   CCTAGAATCTG	2382	GTTTGTTTCCT	1,00	0,22		protein [H.sapiens]
2384   CCTAGAATCTG	2383	AAGACACTGTT	1,00	0,22	Hs.20707	hypothetical protein R31240_1
2385   GTCGGGGAGA   1,00   0,22   Hs.18844   ESTs     2386   ATGTTGTCAAT   1,00   0,22   Hs.1845   MHC class I region ORF     2387   TTTTCCCTCAG   1,00   0,22   Hs.184242   sterol-C5-desaturase (fungal ERG3, delta-5-desaturase) (fungal ERG3, delta-5-desaturase)     2388   ACATTTCATTA   1,00   0,22   Hs.18192   Ser/Arg-related nuclear matrix protein (plenty of pr (plenty of pr (plenty of pr (plenty))   0,22   Hs.180610   Splicing factor proline/glutamine rich (polypyrimidi)     2390   GGTGGAAAAAA   1,00   0,22   Hs.178728   methyl-CpG binding domain protein 3     2391   GGCAGAATAA   1,00   0,22   Hs.163593   ribosomal protein L18a     2392   GGTGCCCGCA   1,00   0,22   Hs.163593   ribosomal protein L18a     2393   TAGCTGAGGCA   1,00   0,22   Hs.159557   karyopherin alpha 2 (RAG cohort 1, importin alpha 1)     2394   CGGATTCAATT   1,00   0,22   Hs.14763   ESTs     2395   AAGAAAGGGAA   1,00   0,22   Hs.14763   ESTs     2396   TCAAAAGGGCA   1,00   0,22   Hs.18738   KIAA0800 gene product     2398   TTGCTGCAGAA   1,00   0,22   Hs.118738   KIAA0800 gene product     2398   TTGCTGCAGAA   1,00   0,22   Hs.110373   ESTs     2400   AGCTTCCGCTT   1,00   0,22   Hs.104311   novel protein with MAM domain     2402   GCCTACCCGAG   15,00   1,89   Hs.23582   tumor-associated calcium signal     2403   GCCCACAGTAG   4,00   0,72   Hs.76122   Splicing factor, arginine/serine-rich 4     2404   ACTGCAGAGCG   4,00   0,72   Hs.72131   insulin-like growth factor binding protein     6   COL01292	2384	CCTAGAATCTG		0,22		
2386   ATGTTGTCAAT   1,00   0,22   Hs.1845   MHC class I region ORF   1,00   0,22   Hs.184242   steroI-C5-desaturase (fungal ERG3, delta-5-desaturase)   delta-5-desaturase (fungal ERG3, delta-5-desaturase)   Ser/Arg-related nuclear matrix protein (plenty of pr (plen	2385	GTCGGGGGAGA		0,22		
2387   TTTTCCCTCAG						
2388   ACATTTCATTA	2387	TTTTCCCTCAG		0,22		sterol-C5-desaturase (fungal ERG3,
1,00	2388	ACATTTCATTA	1,00	0,22	Hs.18192	Ser/Arg-related nuclear matrix protein
Carrier   Carr	2389	TTAGCCAGGGT	1,00	0,22	Hs.180610	splicing factor proline/glutamine rich
2391   GGCAGAATAA	2390	GGTGGAAAAA	1,00	0,22	Hs.178728	
2392   GGTGCCCGGCA   1,00   0,22   Hs.163593   ribosomal protein L18a	2391	GGGCAGAATAA				
TAGCTGAGGCA   1,00   0,22   Hs.159557   karyopherin alpha 2 (RAG cohort 1, importin alpha 1)	2392	GGTGCCCGGCA		0.22	Hs.163593	ribosomal protein L18a
2394 CGGATTCAATT         1,00         0,22 Hs.14763         ESTs           2395 AAGAAAGGGAA         1,00         0,22 Hs.140908         ESTs           2396 TCAAAAGGGCA         1,00         0,22 Hs.125158         ESTs           2397 CTCTCCTTGCC         1,00         0,22 Hs.118738         KIAA0800 gene product           2398 TTGCAGAGGGG         1,00         0,22 Hs.110373         ESTs           2399 TGTCTGCAGAA         1,00         0,22 Hs.107418         ESTs           2400 AGCTTCCGCTT         1,00         0,22 Hs.106529         CGI-65 protein           2401 CTCACACACAC         1,00         0,22 Hs.104311         novel protein with MAM domain           2402 GCCTACCCGAG         15,00         1,89 Hs.23582         tumor-associated calcium signal transducer 2           2403 GCCCACAGTAG         4,00         0,72 Hs.76122         splicing factor, arginine/serine-rich 4           2404 ACTGCAGAGCG         4,00         0,72 Hs.12186         Homo sapiens cDNA FLJ20792 fis, clone COL01292           2405 GGCCCCTCACC         11,00         1,48 Hs.274313         insulin-like growth factor binding protein 6						karyopherin alpha 2 (RAG cohort 1,
2396 TCAAAAGGGCA         1,00         0,22 Hs.125158 ESTs           2397 CTCTCCTTGCC         1,00         0,22 Hs.118738 KIAA0800 gene product           2398 TTGCAGAGGGG         1,00         0,22 Hs.110373 ESTs           2399 TGTCTGCAGAA         1,00         0,22 Hs.107418 ESTs           2400 AGCTTCCGCTT         1,00         0,22 Hs.106529 CGI-65 protein           2401 CTCACACACAC         1,00         0,22 Hs.104311 novel protein with MAM domain           2402 GCCTACCCGAG         15,00         1,89 Hs.23582 tumor-associated calcium signal transducer 2           2403 GCCCACAGTAG         4,00         0,72 Hs.76122 splicing factor, arginine/serine-rich 4           2404 ACTGCAGAGCG         4,00         0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292           2405 GGCCCCTCACC         11,00         1,48 Hs.274313 insulin-like growth factor binding protein 6	2394	CGGATTCAATT	1,00	0,22	Hs.14763	
2396 TCAAAAGGGCA         1,00         0,22 Hs.125158 ESTs           2397 CTCTCCTTGCC         1,00         0,22 Hs.118738 KIAA0800 gene product           2398 TTGCAGAGGGG         1,00         0,22 Hs.110373 ESTs           2399 TGTCTGCAGAA         1,00         0,22 Hs.107418 ESTs           2400 AGCTTCCGCTT         1,00         0,22 Hs.106529 CGI-65 protein           2401 CTCACACACAC         1,00         0,22 Hs.104311 novel protein with MAM domain           2402 GCCTACCCGAG         15,00         1,89 Hs.23582 tumor-associated calcium signal transducer 2           2403 GCCCACAGTAG         4,00         0,72 Hs.76122 splicing factor, arginine/serine-rich 4           2404 ACTGCAGAGCG         4,00         0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292           2405 GGCCCCTCACC         11,00         1,48 Hs.274313 insulin-like growth factor binding protein 6	_					ESTs
2397 CTCTCCTTGCC         1,00         0,22 Hs.118738 KIAA0800 gene product           2398 TTGCAGAGGGG         1,00         0,22 Hs.110373 ESTs           2399 TGTCTGCAGAA         1,00         0,22 Hs.107418 ESTs           2400 AGCTTCCGCTT         1,00         0,22 Hs.106529 CGI-65 protein           2401 CTCACACACAC         1,00         0,22 Hs.104311 novel protein with MAM domain           2402 GCCTACCCGAG         15,00         1,89 Hs.23582 tumor-associated calcium signal transducer 2           2403 GCCCACAGTAG         4,00         0,72 Hs.76122 splicing factor, arginine/serine-rich 4           2404 ACTGCAGAGCG         4,00         0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292           2405 GGCCCCTCACC         11,00         1,48 Hs.274313 insulin-like growth factor binding protein 6						
2398 TTGCAGAGGGG         1,00         0,22 Hs.110373 ESTs           2399 TGTCTGCAGAA         1,00         0,22 Hs.107418 ESTs           2400 AGCTTCCGCTT         1,00         0,22 Hs.106529 CGI-65 protein           2401 CTCACACACAC         1,00         0,22 Hs.104311 novel protein with MAM domain           2402 GCCTACCCGAG         15,00         1,89 Hs.23582 tumor-associated calcium signal transducer 2           2403 GCCCACAGTAG         4,00         0,72 Hs.76122 splicing factor, arginine/serine-rich 4           2404 ACTGCAGAGCG         4,00         0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292           2405 GGCCCCTCACC         11,00         1,48 Hs.274313 insulin-like growth factor binding protein 6						<del></del>
2399 TGTCTGCAGAA         1,00         0,22 Hs.107418 ESTs           2400 AGCTTCCGCTT         1,00         0,22 Hs.106529 CGI-65 protein           2401 CTCACACACAC         1,00         0,22 Hs.104311 novel protein with MAM domain           2402 GCCTACCCGAG         15,00         1,89 Hs.23582 tumor-associated calcium signal transducer 2           2403 GCCCACAGTAG         4,00         0,72 Hs.76122 splicing factor, arginine/serine-rich 4           2404 ACTGCAGAGCG         4,00         0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292           2405 GGCCCCTCACC         11,00         1,48 Hs.274313 insulin-like growth factor binding protein 6						
2400 AGCTTCCGCTT1,000,22 Hs.106529 CGI-65 protein2401 CTCACACACAC1,000,22 Hs.104311 novel protein with MAM domain2402 GCCTACCCGAG15,001,89 Hs.23582 tumor-associated calcium signal transducer 22403 GCCCACAGTAG4,000,72 Hs.76122 splicing factor, arginine/serine-rich 42404 ACTGCAGAGCG4,000,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL012922405 GGCCCCTCACC11,001,48 Hs.274313 insulin-like growth factor binding protein 6						
2401 CTCACACACAC1,000,22 Hs.104311 novel protein with MAM domain2402 GCCTACCCGAG15,001,89 Hs.23582 tumor-associated calcium signal transducer 22403 GCCCACAGTAG4,000,72 Hs.76122 splicing factor, arginine/serine-rich 42404 ACTGCAGAGCG4,000,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL012922405 GGCCCCTCACC11,001,48 Hs.274313 insulin-like growth factor binding protein 6						
2402 GCCTACCCGAG 15,00 1,89 Hs.23582 tumor-associated calcium signal transducer 2  2403 GCCCACAGTAG 4,00 0,72 Hs.76122 splicing factor, arginine/serine-rich 4  2404 ACTGCAGAGCG 4,00 0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292  2405 GGCCCTCACC 11,00 1,48 Hs.274313 insulin-like growth factor binding protein 6						
2403 GCCCACAGTAG 4,00 0,72 Hs.76122 splicing factor, arginine/serine-rich 4 2404 ACTGCAGAGCG 4,00 0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292 2405 GGCCCCTCACC 11,00 1,48 Hs.274313 insulin-like growth factor binding protein 6						tumor-associated calcium signal
2404 ACTGCAGAGCG 4,00 0,72 Hs.12186 Homo sapiens cDNA FLJ20792 fis, clone COL01292 2405 GGCCCTCACC 11,00 1,48 Hs.274313 insulin-like growth factor binding protein 6	2403	GCCCACAGTAG	4.00	0.72	Hs.76122	
2405 GGCCCCTCACC 11,00 1,48 Hs.274313 insulin-like growth factor binding protein 6						Homo sapiens cDNA FLJ20792 fis, clone
	2405	GGCCCCTCACC	11,00	1,48	Hs.274313	insulin-like growth factor binding protein
	2406	AACAGATATTG	3.00	0.58	Hs.190161	<u> </u>

2407	GTGGCAGACGC	3,00			ESTs, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ
2408	GAAAATAAAGT	3,00	0,58	Hs.111334	ferritin, light polypeptide
2409	TTAGTGTCGTA	11,00	1,47	Hs.74649	cytochrome c oxidase subunit VIc
2410	CTTCCTTGCCT	18,00	2,10	Hs.2785	keratin 17
2411	TTCATAGCTGC	8,00	1,16	Hs.155481	cartilage associated protein
2412	CCCTAGGTTGG	13,00	1,64	Hs.3989	plexin B2
2413	AAGATATCAGT	2,00		Hs.74122	caspase 4, apoptosis-related cysteine protease
2414	TTTGTTGAATG	2,00	0,42	Hs.44856	ESTs, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ
2415	CAACGTCCTGA	2,00	0,42	Hs.29068	ESTs
2416	GAATTGACGCC	2,00	0,42	Hs.21851	Homo sapiens mRNA; cDNA DKFZp586J2118 (from clone DK
2417	AACCCATTGTG	2,00	0,42	Hs.184326	CDC10 (cell division cycle 10, S. cerevisiae, homolo
2418	ACCACTTCCTC	2,00			Human clone CE29 8.1 (CAC)n/(GTG)n repeat-containing
2419	GCCCATCGTAC	2,00	0,42		ESTs, Weakly similar to testicular tektin B1-like pr
2420	CCTACTCCCAG	2,00	0,42	Hs.10071	seven transmembrane protein TM7SF3
2421	GCCTGGTGACC	5,00	0,83	Hs.180224	death-associated protein 6
2422	ATGGCAACAGA	13,00	1,62		integrin, alpha 5 (fibronectin receptor, alpha polyp
	CAGGCCTGGCC	8,00	1,15	Hs.74649	cytochrome c oxidase subunit VIc
	GTGAGACCCCG	8,00	1,15		ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
2425	GAGCTGGTGAA	3,00	0,57		ESTs, Weakly similar to myotonic dystrophy kinase [H
2426	TCTGTTGAGTG	3,00	0,57		ESTs
2427	CTCAACCCCCC	15,00	1,78		low density lipoprotein-related protein 1 (alpha-2-m
2428	ATCACTAAAGA	4,00	0,70	Hs.75888	phosphogluconate dehydrogenase
-	CCCCTGCACTC	4,00	0,70	Hs.235213	ESTs
	CCCTCTGTCAG	4,00			Homo sapiens mRNA; cDNA DKFZp761E0711 (from clone DK
2431	CACTGCCTTTG	4,00	0,70	Hs.106019	protein phosphatase 1, regulatory subunit 10
2432	AAGATCAAGAT	10,00	1,32		actin, alpha 1, skeletal muscle
2433	GCGAGACCCTG	8,00	1,13		Homo sapiens mRNA; cDNA DKFZp434A1014 (from clone DK
2434	TTTGCCTGGAT	2,00	0,41	Hs.95260	Autosomal Highly Conserved Protein
2435	GTCCCAACACA	2,00		Hs.8961	ESTs
2436	TTGGGGTGCCT	2,00			hydroxysteroid (17-beta) dehydrogenase
	AAAAGGCACTT	2,00			actin related protein 2/3 complex, subunit 5 (16 kD)
2438	TCTTACGCGTT	2,00	0,41	Hs.7905	SH3 and PX domain-containing protein SH3PX1

<b>-</b> (	1470040	0.00		05:55	. 511
2439	ATGCAGTTCAA	2,00	0,41	Hs.65135	Homo sapiens mRNA; cDNA
2440	AT007040400	0.00	0.44	11- 0405	DKFZp434E0121 (from clone DK
2440	ATGCTGAGAGG	2,00	0,41	Hs.6185	ESTs, Weakly similar to
0444	04704040700	0.00	0.44	11- 50700	BcDNA.GH12174 [D.melanogaste
	CATCACACTCC	2,00			lymphocyte-specific protein 1
2442	тестеттесте	2,00	0,41	Hs.55923	ESTs, Highly similar to lin-10 protein
2/12					homolog [R.no
-	ATTGTTTCTTG	2,00			KIAA0867 protein
	AGGGAGACCTG	2,00			jumonji (mouse) homolog
	CTACAAAAAGA_	2,00			mitochondrial carrier homolog 2
	TGTGTTAAGAG	2,00			hypothetical protein FLJ20061
	CCACTATACTC	2,00		Hs.269404	
2448	GAATGTCCTTT	2,00	0,41	Hs.172674	nuclear factor of activated T-cells, cytoplasmic 3
2449	GAACGACCTAG	2,00	0,41	Hs.15783	Homo sapiens mRNA; cDNA
					DKFZp434P1115 (from clone DK
2450	AAGAAGGCAAG	2,00	0,41	Hs.155402	D site of albumin promoter (albumin D-
					box) binding p
2451	ATCTATTGAAA	2,00	0,41	Hs.15386	ESTs
2452	TCCTATGCAGA	2,00	0,41	Hs.108924	DKFZP586P1422 protein
2453	GAGACCCTGGA	3,00		Hs.8088	similar to S. cerevisiae Sec6p and R.
			·		norvegicus rse
2454	TAGCTCGATCC	3,00	0,56	Hs.58372	ESTs
	TTGTCCAGGCT	7,00			ATP-dependent RNA helicase
	ACCCTTTAACA	11,00			major histocompatibility complex, class I,
2457	TTTTTGTATTA	4,00	0,69	Hs.179526	upregulated by 1,25-dihydroxyvitamin D-
2450	CCACATCACCA	5.00	0.04	11- 02440	UIAA0050 - makain
	GGAGATGAGGA	5,00			KIAA0252 protein
	AGCCTGGGAGG	5,00			CGI-43 protein
	TCCCTGGGGGT	1,00			ESTs
	ATCAAAGGTTT	1,00	_	Hs.94795	Homo sapiens mRNA; cDNA DKFZp564O222 (from clone DKF
2462	TAACTTTATAA	1,00			KIAA0579 protein
	TTGTCCTGACC	1,00	0,20	Hs.80181	ESTs
2464	AACTATGCCAG	1,00	0,20	Hs.7905	SH3 and PX domain-containing protein SH3PX1
2465	CTGCTGTAATA	1,00	0,20	Hs.77508	glutamate dehydrogenase 1
	GGGTCCTCTCC	1,00		Hs.76064	ribosomal protein L27a
	AGGGCAGTGCC	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	TCACCGTGGAC	1,00		Hs.73800	selectin P (granule membrane protein
		.,,,,,	-,-0		140kD, antigen
2469	AACCCGGTAGG	1,00	0.20	Hs.69707	H.sapiens HCG II mRNA
	ACACTGATTAA	1,00		Hs.59838	hypothetical protein FLJ10808
	GGGTTGGGGTC	1,00		Hs.58014	G protein-coupled receptor, family C,
		.,,,,,	-,-0		group 5, membe
2472	GCATTGTGGTG	1,00	0.20	Hs.5534	ESTs
	GATTATGTAAT	1,00		Hs.5056	CGI-41 protein
	TATGAGCAGAA	1,00		Hs.5010	Homo sapiens clone 24672 mRNA
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0.475	T4440004040	4.00	0.00	11- 4000	sequence
	TAAAGGCACAG	1,00		Hs.4980	LIM domain binding 2
	AGCAAAACTGG	1,00		Hs.46489	ESTs
24//	TGGAACTGAGT	1,00	0,20	Hs.43899	Homo sapiens mRNA; cDNA
0.470	OTOATTOAAOT	4.00	0.00	11: 00040	DKFZp434C1714 (from clone DK
	CTGATTCAACT	1,00		Hs.30818	ESTs
24/9	GGTGAGCTACT	1,00	0,20	Hs.29874	ESTs, Weakly similar to A49656
0.400	00011111	4.00			estrogen-responsive f
	CCCAAAAAAA	1,00			JM4 protein
2481	GGTTATTGTAT	1,00	0,20	Hs.286184	Homo sapiens mRNA; cDNA
0.400	CACAACTOTAC	4.00	0.00	11 000100	DKFZp566C114 (from clone DKF
	GAGAACTGTAG	1,00		Hs.286169	
	TCTGAAAAAA	1,00		Hs.285112	
2484	GGGACGGGGTG	1,00	0,20	Hs.284161	Homo sapiens mRNA for KIAA1532
2.425		4.55			protein, partial cds
$\overline{}$	TGTAGGGGTTA	1,00			polymerase (DNA directed), alpha
2486	AAGCTGGAGGC	1,00	0,20	Hs.265327	Human DNA sequence from clone RP4-
					756G23 on chromoso
	TACAAAGCATA	1,00			HIRA-interacting protein 3
	AATTAAAAAAA	1,00		Hs.257515	
	GAACACCGTCC	1,00			hypothetical protein DKFZp761G1923
	GCAAAACTGTC	1,00		Hs.252721	
	GCCGCCCTACG	1,00			coronin, actin-binding protein, 2B
2492	GGGCCTTGGAG	1,00	0,20		mitogen-activated protein kinase-
					activated protein k
2493	GGAAGACCAGA	1,00	0,20		ESTs, Weakly similar to cDNA EST
					yk282b7.5 comes fro
2494	GTGGCTGACGC	1,00	0,20		ESTs, Weakly similar to ALU4_HUMAN
2 12 2					ALU SUBFAMILY SB2
2495	TCAAGTTTAAA	1,00	0,20		myeloid/lymphoid or mixed-lineage
0.400					leukemia (trithora
2496	AAACTATCACA	1,00	0,20		solute carrier family 7 (cationic amino
0.40=	0007407070				acid transpo
2497	GGGTAGTGTCA	1,00	0,20	Hs.183738	FERM, RhoGEF (ARHGEF) and
0.400	TTT-1-1-0-0	4.00			pleckstrin domain protein
	TTTTATTATCC	1,00			actinin, alpha 4
2499	GCCTTTCCTCA	1,00	0,20		conserved gene amplified in
0500	CATTTACTOTA	4 00	0.00		osteosarcoma
	CATTTACTCTA	1,00	0,20	Hs.17109	integral membrane protein 2A
2501	CTGCCATCTTT	1,00	0,20		Homo sapiens mRNA; cDNA
0500	TOTTOTTOTA	- 4 3 3			DKFZp586K1924 (from clone DK
	TCTTCTTTGTA	1,00		Hs.160711	
	GGCAAATGAAG	1,00			PPAR binding protein
_	GTGGTTTGTTG	1,00			KIAA0740 gene product
	TTAGGGCCCAG	1,00		Hs.137313	
	GCCCTTCTCAG	1,00			beta V spectrin
	CTTGCTGAAGA	1,00		Hs.13619	
	AAGTAAGTCTA	1,00			CGI-43 protein
2509	GCACCAGGAGC	1,00	0,20	Hs.110533	Homo sapiens mRNA; cDNA

DKFZp761H172 (from clone DKF	
2511   TTTGAATCAGT	
2512   TTTTGTGTTGG	
2513   CAAACCTTTAA	
2514   CTAAAAAAAA   31,00   2,97   Hs.23740   KIAA1598 protein   2515   CCAGTAATCCC   7,00   1,01   Hs.237078   ESTs   2516   CCCATAATCCC   7,00   1,01   Hs.111256   arachidonate 15-lipoxygenase, seco type   2517   ATGGCACCACT   5,00   0,79   Hs.161554   hypothetical protein FLJ20159   2518   CAGCTGGCCAT   9,00   1,19   Hs.79732   fibulin 1   2519   CTGCTGCACTC   9,00   1,19   Hs.193974   ESTs, Weakly similar to ALU1_HUMALU SUBFAMILY J S   2520   AATGGCATTGA   4,00   0,68   Hs.31431   ESTs   2521   CTCCCCTGCCC   9,00   1,18   Hs.82422   capping protein (actin filament), gels like   2522   ATGTCCAATTT   3,00   0,55   Hs.247309   succinate-CoA ligase, GDP-forming, subunit   2523   CTACAATTTC   3,00   0,55   Hs.24307   f-box and WD-40 domain protein 3   2524   CTGTAAAACAA   3,00   0,55   Hs.180686   ubiquitin protein ligase E3A (human papilloma virus   2525   TGAGGGATGGA   3,00   0,55   Hs.172740   microtubule-associated protein, RP/I family, member   2526   AGGCAGAGGTT   3,00   0,55   Hs.164129   ESTs, Weakly similar to ALU6_HUMALU SUBFAMILY SP   hypothetical protein HDCMC04P   2528   ATGAAACCTG   2,00   0,40   Hs.78353   SFRS protein kinase 2   2530   CTGAAGGCTGA   2,00   0,40   Hs.78146   platelet/endothelial cell adhesion molecule (CD31 an   2531   TGACAATTTG   2,00   0,40   Hs.78146   platelet/endothelial cell adhesion molecule (CD31 an   2532   TGCATCTGTGC   2,00   0,40   Hs.78146   platelet/endothelial cell adhesion molecule (CD31 an   2534   AAGGGGCCTTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1   2535   CCACTGTACTT   2,00   0,40   Hs.26208   co	
2515   CCAGTAATCCC   7,00   1,01   Hs.237078   ESTs	
2516   CCCATAATCCC   7,00   1,01   Hs.111256   arachidonate 15-lipoxygenase, secon type	
type	
2517   ATGGCACCACT   5,00   0,79   Hs.161554   hypothetical protein FLJ20159	חם
2518   CAGCTGGCCAT   9,00   1,19   Hs.79732   fibulin 1	
2519 CTGCTGCACTC   9,00   1,19   Hs.193974   ESTs, Weakly similar to ALU1_HUM   ALU SUBFAMILY J S	
ALU SUBFAMILY J S	A A A
2521   CTCCCTGCCC   9,00   1,18   Hs.82422   capping protein (actin filament), gels like     2522   ATGTCCAATTT   3,00   0,55   Hs.247309   succinate-CoA ligase, GDP-forming, subunit     2523   CTACAATTTC   3,00   0,55   Hs.24307   f-box and WD-40 domain protein 3     2524   CTGTAAAACAA   3,00   0,55   Hs.180686   ubiquitin protein ligase E3A (human papilloma virus     2525   TGAGGGATGGA   3,00   0,55   Hs.172740   microtubule-associated protein, RP/I family, member     2526   AGGCAGAGGTT   3,00   0,55   Hs.164129   ESTs, Weakly similar to ALU6_HUM     ALU SUBFAMILY SP     2527   GAAAATATCAA   3,00   0,55   Hs.15423   hypothetical protein HDCMC04P     2528   ATGAAACCCTG   27,00   2,59   Hs.187991   DKFZP564A122 protein     2529   GTCTTCGAAGT   2,00   0,40   Hs.78353   SFRS protein kinase 2     2530   CTGAAGGCTGA   2,00   0,40   Hs.78146   platelet/endothelial cell adhesion molecule (CD31 an     2531   TGACAATTTTG   2,00   0,40   Hs.58589   glycogenin 2     2533   ACCGAAACTTG   2,00   0,40   Hs.38041   pyridoxal (pyridoxine, vitamin B6) kir     2534   AAGGGGCCTTT   2,00   0,40   Hs.26208   collagen, type XVI, alpha 1     2535   CCACTGTACTT   2,00   0,40   Hs.183475   Homo sapiens clone 25061 mRNA     sequence   Sapina   Subunit   subun	AN 
Sike	
Subunit   Subu	olin-
2523 CTACAATTITC         3,00         0,55 Hs.24307 f-box and WD-40 domain protein 3           2524 CTGTAAAACAA         3,00         0,55 Hs.180686 ubiquitin protein ligase E3A (human papilloma virus)           2525 TGAGGGATGGA         3,00         0,55 Hs.172740 microtubule-associated protein, RP/I family, member           2526 AGGCAGAGGTT         3,00         0,55 Hs.164129 ESTs, Weakly similar to ALU6_HUM ALU SUBFAMILY SP           2527 GAAAATATCAA         3,00         0,55 Hs.15423 hypothetical protein HDCMC04P           2528 ATGAAACCCTG         27,00         2,59 Hs.187991 DKFZP564A122 protein           2529 GTCTTCGAAGT         2,00         0,40 Hs.78353 SFRS protein kinase 2           2530 CTGAAGGCTGA         2,00         0,40 Hs.78146 platelet/endothelial cell adhesion molecule (CD31 an           2531 TGACAATTITG         2,00         0,40 Hs.75912 KIAA0257 protein           2532 TGCATCTGTGC         2,00         0,40 Hs.58589 glycogenin 2           2533 ACCGAAACTTG         2,00         0,40 Hs.38041 pyridoxal (pyridoxine, vitamin B6) kir           2534 AAGGGGCCTTT         2,00         0,40 Hs.26208 collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	beta
2524 CTGTAAAACAA 3,00 0,55 Hs.180686 ubiquitin protein ligase E3A (human papilloma virus) 2525 TGAGGGATGGA 3,00 0,55 Hs.172740 microtubule-associated protein, RP/II family, member 2526 AGGCAGAGGTT 3,00 0,55 Hs.164129 ESTs, Weakly similar to ALU6_HUMALU SUBFAMILY SP 2527 GAAAATATCAA 3,00 0,55 Hs.15423 hypothetical protein HDCMC04P 2528 ATGAAACCCTG 27,00 2,59 Hs.187991 DKFZP564A122 protein 2529 GTCTTCGAAGT 2,00 0,40 Hs.78353 SFRS protein kinase 2 2530 CTGAAGGCTGA 2,00 0,40 Hs.78146 platelet/endothelial cell adhesion molecule (CD31 an 2531 TGACAATTITG 2,00 0,40 Hs.75912 KIAA0257 protein 2532 TGCATCTGTGC 2,00 0,40 Hs.58589 glycogenin 2 2533 ACCGAAACTTG 2,00 0,40 Hs.38041 pyridoxal (pyridoxine, vitamin B6) kir 2534 AAGGGGCCTTT 2,00 0,40 Hs.26208 collagen, type XVI, alpha 1 2535 CCACTGTACTT 2,00 0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	
papilloma virus	
2525         TGAGGGATGGA         3,00         0,55         Hs.172740         microtubule-associated protein, RP/II family, member           2526         AGGCAGAGGTT         3,00         0,55         Hs.164129         ESTs, Weakly similar to ALU6_HUM ALU SUBFAMILY SP           2527         GAAAATATCAA         3,00         0,55         Hs.15423         hypothetical protein HDCMC04P           2528         ATGAAACCCTG         27,00         2,59         Hs.187991         DKFZP564A122 protein           2529         GTCTTCGAAGT         2,00         0,40         Hs.78353         SFRS protein kinase 2           2530         CTGAAGGCTGA         2,00         0,40         Hs.78146         platelet/endothelial cell adhesion molecule (CD31 an           2531         TGACAATTTTG         2,00         0,40         Hs.75912         KIAA0257 protein           2532         TGCATCTGTGC         2,00         0,40         Hs.58589         glycogenin 2           2533         ACCGAAACTTG         2,00         0,40         Hs.38041         pyridoxal (pyridoxine, vitamin B6) kir           2534         AAGGGGCCTTT         2,00         0,40         Hs.26208         collagen, type XVI, alpha 1           2535         CCACTGTACTT         2,00         0,40         Hs.183475         H	
Second	В
ALU SUBFAMILY SP	
2528 ATGAAACCCTG         27,00         2,59 Hs.187991 DKFZP564A122 protein           2529 GTCTTCGAAGT         2,00         0,40 Hs.78353 SFRS protein kinase 2           2530 CTGAAGGCTGA         2,00         0,40 Hs.78146 platelet/endothelial cell adhesion molecule (CD31 an           2531 TGACAATTTTG         2,00         0,40 Hs.75912 KIAA0257 protein           2532 TGCATCTGTGC         2,00         0,40 Hs.58589 glycogenin 2           2533 ACCGAAACTTG         2,00         0,40 Hs.38041 pyridoxal (pyridoxine, vitamin B6) kir           2534 AAGGGGCCTTT         2,00         0,40 Hs.26208 collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	AN
2528 ATGAAACCCTG         27,00         2,59 Hs.187991 DKFZP564A122 protein           2529 GTCTTCGAAGT         2,00         0,40 Hs.78353 SFRS protein kinase 2           2530 CTGAAGGCTGA         2,00         0,40 Hs.78146 platelet/endothelial cell adhesion molecule (CD31 an           2531 TGACAATTTTG         2,00         0,40 Hs.75912 KIAA0257 protein           2532 TGCATCTGTGC         2,00         0,40 Hs.58589 glycogenin 2           2533 ACCGAAACTTG         2,00         0,40 Hs.38041 pyridoxal (pyridoxine, vitamin B6) kir           2534 AAGGGGCCTTT         2,00         0,40 Hs.26208 collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	
2530 CTGAAGGCTGA         2,00         0,40 Hs.78146         platelet/endothelial cell adhesion molecule (CD31 an           2531 TGACAATTTTG         2,00         0,40 Hs.75912         KIAA0257 protein           2532 TGCATCTGTGC         2,00         0,40 Hs.58589         glycogenin 2           2533 ACCGAAACTTG         2,00         0,40 Hs.38041         pyridoxal (pyridoxine, vitamin B6) kir           2534 AAGGGGCCTTT         2,00         0,40 Hs.26208         collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475         Homo sapiens clone 25061 mRNA sequence	
molecule (CD31 an   2531 TGACAATTITG   2,00   0,40 Hs.75912   KIAA0257 protein   2532 TGCATCTGTGC   2,00   0,40 Hs.58589   glycogenin 2   2533 ACCGAAACTTG   2,00   0,40 Hs.38041   pyridoxal (pyridoxine, vitamin B6) kir   2534 AAGGGGCCTTT   2,00   0,40 Hs.26208   collagen, type XVI, alpha 1   2535 CCACTGTACTT   2,00   0,40 Hs.183475   Homo sapiens clone 25061 mRNA sequence	
2531 TGACAATTTG         2,00         0,40 Hs.75912         KIAA0257 protein           2532 TGCATCTGTGC         2,00         0,40 Hs.58589         glycogenin 2           2533 ACCGAAACTTG         2,00         0,40 Hs.38041         pyridoxal (pyridoxine, vitamin B6) kir           2534 AAGGGGCCTTT         2,00         0,40 Hs.26208         collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475         Homo sapiens clone 25061 mRNA sequence	
2532 TGCATCTGTGC         2,00         0,40 Hs.58589         glycogenin 2           2533 ACCGAAACTTG         2,00         0,40 Hs.38041         pyridoxal (pyridoxine, vitamin B6) kir           2534 AAGGGGCCTTT         2,00         0,40 Hs.26208         collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475         Homo sapiens clone 25061 mRNA sequence	
2533 ACCGAAACTTG 2,00 0,40 Hs.38041 pyridoxal (pyridoxine, vitamin B6) kir 2534 AAGGGCCTTT 2,00 0,40 Hs.26208 collagen, type XVI, alpha 1 2535 CCACTGTACTT 2,00 0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	
2534 AAGGGGCCTTT         2,00         0,40 Hs.26208 collagen, type XVI, alpha 1           2535 CCACTGTACTT         2,00         0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	
2535 CCACTGTACTT 2,00 0,40 Hs.183475 Homo sapiens clone 25061 mRNA sequence	ase
sequence	
2536 ATGCTTTTCAC 2,00 0,40 Hs.181357 laminin receptor 1 (67kD, ribosomal	
protein SA)	
2537 GGGTCAGGAGA 2,00 0,40 Hs.181271 CGI-120 protein	
2538 CACATCCTTAC 2,00 0,40 Hs.173717 phosphatidic acid phosphatase type	2B
2539 AATATTAAAAA 2,00 0,40 Hs.168103 prp28, U5 snRNP 100 kd protein	$\neg$
2540 GGCCCCCTCC 2,00 0,40 Hs.155979 KIAA0295 protein	
2541 TCTAAAAAAA 2,00 0,40 Hs.13328 ESTs	
2542 CGCCTATAATC 6,00 0,88 Hs.194110 Homo sapiens mRNA; cDNA	
DKFZp434C0814 (from clone DK	ı
2543 AAGGATGCCAA 10,00 1,25 Hs.169946 GATA-binding protein 3	$\neg$
2544 GTCCCTGCCTT 4,00 0,67 Hs.279837 glutathione S-transferase M2 (musc	э)
2545 CACCCAATGGG 4,00 0,67 Hs.110121 SEC7 homolog	
2546 CAGGAGGAAAG 5,00 0,77 Hs.177425 KIAA0964 protein	

2547	TGGGCCCGTGT	5,00	0,77	Hs.11607	ESTs
	TGTGACCTCTC	5,00	0,77	Hs.108973	dolichyl-phosphate mannosyltransferase
			Ť		polypeptide 2
2549	GGAGGCGGAGG	6,00	0,87	Hs.15562	hypothetical protein FLJ10871
	GGACAGCTCAG	3,00			ESTs, Weakly similar to hypothetical
	•				protein [H.sapi
2551	GTTACCAGTTT	3,00	0,54	Hs.28264	Homo sapiens mRNA; cDNA
					DKFZp564L0822 (from clone DK
2552	CAACTTAAGTG	3,00	0,54	Hs.16492	DKFZP564G2022 protein
2553	CCAGTGCACTC	10,00	1,23	Hs.254856	EST
2554	TGCACACACAC	4,00			beta-catenin-interacting protein ICAT
2555	CCTCTAATCCC	4,00	0,66	Hs.236150	ESTs, Weakly similar to AF090942_1 PRO0657 [H.sapien
2556	AAGCGCTCTCG	6,00	0,86	Hs.168913	serine/threonine kinase 24 (Ste20, yeast homolog)
2557	CTTGTGTGTAG	6,00	0,86	Hs.158203	actin binding LIM protein 1
2558	GAAATCTGTCC	2,00	0,39	Hs.8710	ESTs
2559	CAAGGAGATCT	2,00	0,39	Hs.7471	ESTs, Weakly similar to homology with GTP binding pr
2560	CAAAATACTGC	2,00	0,39	Hs.6557	ESTs
2561	AGATAGCATTA	2,00		Hs.6241	phosphoinositide-3-kinase, regulatory subunit, polyp
2562	AGAATGTACGG	2,00	0,39	Hs.57973	hypothetical protein
	CCACGCACTGT	2,00	0,39	Hs.48778	Homo sapiens mRNA; cDNA DKFZp586O0221 (from clone DK
2564	CACCCCTCAGG	2,00	0,39	Hs.26369	hypothetical protein FLJ20287
2565	ATTACACCACT	2,00	0,39	Hs.254565	EST, Weakly similar to A46010 X-linked retinopathy p
2566	AACCCAGAAGG	2,00	0,39	Hs.229985	EST, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB S
2567	GTGGTGTATGC	2,00	0,39	Hs.229244	
2568	AACTATAAACG	2,00	0,39	Hs.22393	density-regulated protein
2569	AAGTTTATAGA	2,00	0,39	Hs.206097	oncogene TC21
2570	GTGAAGTTGCG	2,00	0,39	Hs.123178	translocase of inner mitochondrial membrane 44 (yeas
2571	CCCTCCTGCTC	3,00	0,53	Hs.96731	huntingtin interacting protein-1-related
2572	AACCCAAACTC	3,00		Hs.11184	hypothetical protein FLJ20419
2573	GTTGGGACATC	4,00		Hs.9663	programmed cell death 6-interacting protein
2574	CCGCTGATCCA	4,00	0.65	Hs.184161	exostoses (multiple) 1
	GGATACAACCT	4,00			RNA binding motif protein 6
	ATTTAAAAAAA	4,00	0,65	Hs.1139	cold shock domain protein A
	CATTTAAAAAA	1,00		Hs.9443	zinc finger protein 202
	TATACCTGTGT	1,00		Hs.92200	KIAA0480 gene product
2579	GTGAATGACAA	1,00		Hs.8834	ring finger protein 3
	ССССТССССТС	1,00		Hs.83429	tumor necrosis factor (ligand) superfamily, member 1
2581	CAATTTATATC	1,00	0.19	Hs.8294	KIAA0196 gene product
	ACAGATTCAAT	1,00		Hs.8248	NADH dehydrogenase (ubiquinone) Fe-S

		·			
					protein 1 (75kD
	AGTGTATCACA	1,00		Hs.79844	DKFZP564M1416 protein
2584	GAGGGAGTTGG	1,00	0,19	Hs.77810	nuclear factor of activated T-cells, cytoplasmic 4
2595	TAACATCCCTG	1,00	0.10	Hs.75400	KIAA0280 protein
	GGGAGCTTGTA	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	TAGATGTGATT	1,00		Hs.62112	zinc finger protein 207
	TATAGATCGTC	1,00		Hs.59757	zinc finger protein 281
-	AAGTTTTGATT	1,00		Hs.58382	hypothetical protein FLJ11101
2590	CCAATAGGGCC	1,00	0,19	Hs.5555	ESTs, Weakly similar to TERA_HUMAN TRANSITIONAL ENDO
2591	TCGCACACTTT	1,00	0,19	Hs.48998	fibronectin leucine rich transmembrane protein 2
2592	AAAGGTTGCAG	1,00	0,19	Hs.47367	Homo sapiens mRNA; cDNA DKFZp4340031 (from cione DKF
2593	AGCTTTCCAAT	1,00	0.19	Hs.45109	ESTs
	TAGCTTCCTTA	1,00			homeo box C10
	CTGTTACCTTC	1,00			ESTs
	CTTATGTATTA	1,00			ESTs
	TCAACACAGTT	1,00		Hs.285874	
	TATTTCGTAAA	1,00		Hs.285533	
	TACCTGTAATC	1,00		Hs.284555	
	AATATGTGTAC	1,00			katanin p60 (ATPase-containing) subunit A 1
2601	TGACCAGGCCC	1,00	0,19	Hs.27536	ESTs, Weakly similar to unnamed protein product [H.s
2602	AAATGGATGCA	1,00	0.19	Hs.271943	tetraspanin TM4-B
2603	GTGAACCCACG	1,00		Hs.270791	
2604	TCCTATCCAGG	1,00			protein tyrosine phosphatase, receptor type, A
2605	TATATATGGGG	1,00	0.19	Hs.25426	ESTs
	GCAGGTACTGA	1,00			solute carrier family 31 (copper transporters), memb
2607	GCGGCGTGTGC	1,00	0,19	Hs.237040	ESTs, Weakly similar to AF119917_17 PRO1847 [H.sapie
2608	TTATTTCTTCT	1,00	0,19	Hs.22246	ESTs
2609	GTGGCGCACAT	1,00	0,19	Hs.210979	EST
2610	TGTTTCATTCT	1,00		Hs.2055	ubiquitin-activating enzyme E1 (A1S9T and BN75 tempe
2611	TTTAAATAGCC	1,00	0,19	Hs.182965	Kruppel-like factor 4 (gut)
	AGAAATAAATG	1,00		Hs.178356	
	TAAAGCACTTA	1,00		Hs.172806	
	GAAATAGGAAG	1,00			KIAA0440 protein
	GATCATACTCC	1,00			Kreisler (mouse) maf-related leucine zipper homolog
2616	CATAATGTCAT	1,00	0 10	Hs.169152	
	GTGTTTTGTT	1,00			ets variant gene 6 (TEL oncogene)
	AGAATTTAAAA	1,00		Hs.158867	
	CAAACATTCAA	1,00			SH3 domain binding glutamic acid-rich
2013	ON TON	1,00	<u> </u>	1 13. 14300	or is domain binding glutainic acid-nch

			_		protein like
2000	TOOTTTOOATT	1 00	0.40	U- 400050	
	TCCTTTGGATT	1,00		Hs.128659	
	TTCTTTCCTGA	1,00		Hs.127453	
2622	AAAAAGGTGGA	1,00	0,19		ESTs, Moderately similar to B43284 zinc
0000	00717000100	4 00	0.40		finger prote
	GGTATGGCAGG	1,00		Hs.116998	
	AATGGAAGGTG	1,00			hypothetical protein FLJ11264
	GGGCTTTCTG	1,00	0,19	Hs.110480	DC12 protein
2626	AAAATAAACTT	1,00			phosphatase and tensin homolog (mutated in multiple
	TAATATGAGCA	1,00		Hs.105642	
2628	ATATACTGTAG	1,00			hypothetical protein FLJ10697
2629	CGCACCGGGTA	1,00			Homo sapiens napsin 2 precursor, mRNA, partial seque
2630	GGAGGTGGGGC	23,00		Hs.180577	
2631	GGAAATGTCAA	17,00			matrix metalloproteinase 2 (gelatinase A, 72kD gelat
2632	TGTTCTGGAGA	22,00	2,05	Hs.74471	gap junction protein, alpha 1, 43kD (connexin 43)
2633	TACTTGGGAGG	5,00	0,75		LIM protein (similar to rat protein kinase C-binding
2634	GCTGTTCAGAA	4,00	0,64		PRO1073 protein
	CTGTCTGTGGC	4,00			hypothetical protein FLJ10209
	GCACCTAGTGC	4,00			zinc finger protein 173
	GAGCCTGTAAA	3,00			ESTs
_	AGCGAGAGAGG	3,00			ESTs, Weakly similar to bromodomain
		-,	0,00		containing prote
2639	AGTAGCTTGAA	3,00	0.52	Hs.154248	KIAA0549 protein
	CTGTCAGCGGC	7,00		Hs.7247	ESTs, Weakly similar to TIE1_HUMAN TYROSINE-PROTEIN
2641	GCCCAGCGGCC	7,00	0.92	Hs.194385	hypothetical protein FLJ20234
	ATGGTGCACAC	2,00		Hs.87521	ESTs, Weakly similar to ORF2
		_,	-,		consensus sequence enco
2643	ATGTCGTGGTC	2,00	0,38	Hs.6900	ring finger protein 13
	GCCCTTCCCCT	2,00	0,38	Hs.63667	transcriptional adaptor 3 (ADA3, yeast homolog)-like
2645	TGAGTTTTACA	2,00	0,38	Hs.58373	ESTs
	CGGGGCGCGCA	2,00	0,38	Hs.4299	ESTs
2647	AGGAAGCTGAG	2,00	0,38	Hs.3041	uracil-DNA glycosylase 2
2648	CACCTGTAGTT	2,00	0,38	Hs.271053	ESTs, Weakly similar to A46010 X-linked retinopathy
2649	CCCTGGGCTTC	2,00	0,38	Hs.226770	DKFZP566C0424 protein
	TAAGGCCTTTC	2,00			leukemia associated gene 1, candidate tumor suppress
2651	AAATCAGGAAC	2,00	0,38	Hs.180549	ESTs, Highly similar to R26660_1, partial CDS [H.sap
2652	GGAAACCCCAG	2,00	0,38	Hs.174044	dishevelled 3 (homologous to Drosophila dsh)
2653	ACGCACATTAT	2,00	0,38	Hs.156007	Down syndrome critical region gene 1-

		· · ·		<del></del>	list- 4
0054	OTTOTOTO	0.00	0.00	11- 450557	like 1
2654	CTTCTCTTGAG	2,00	0,38	HS.15055/	basic transcription element binding protein 1
2655	TTCCTCCAAAA	2,00		Hs.125743	
2656	GATGAGGAGAC	46,00	3,46	Hs.179573	collagen, type I, alpha 2
2657	CTGTGAGACCT	11,00	1,22	Hs.160741	Fc fragment of IgG, receptor, transporter, alpha
2658	CCACACAAGCA	7,00	0.91	Hs.102737	goliath protein
	TTGGCCAGGAT	31,00			Homo sapiens cDNA FLJ10952 fis, clone PLACE1000374
2660	CTCCCTCTGCC	11,00	1,21	Hs.194534	vesicle-associated membrane protein 2 (synaptobrevin
2661	GAATAAATGTT	6,00	0,82	Hs.8762	FK506-binding protein 9 (63 kD)
	ACCAGCTGTCC	3,00		Hs.84153	dynamitin (dynactin complex 50 kD subunit)
	CTTCTATGTAG	3,00		Hs.77225	ADP-ribosyltransferase (NAD+; poly (ADP-ribose) poly
	GTTTCAGTTAC	3,00		Hs.7016	RAB7, member RAS oncogene family
2665	GGCCAGTAACA	3,00		Hs.69559	KIAA1096 protein
2666	GGAGGTGGGAG	3,00	0,51	Hs.105097	thymidine kinase 1, soluble
2667	TGCAATATTTC	4,00	0,62	Hs.94970	KIAA0306 protein
2668	CAGACTTTTGG	5,00	0,72	Hs.63348	DKFZP586M121 protein
2669	GTGGTGCACGC	28,00	2,27	Hs.181318	ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
2670	GACCAGGCCCT	22,00	1,90		tropomyosin 2 (beta)
2671	CGGCAGAGCTG	4,00		Hs.9610	purinergic receptor P2X, ligand-gated ion channel, 4
2672	AACCCAAAAAG	2,00	0,37	Hs.92323	FXYD domain-containing ion transport regulator 3
2673	TTGCCTTGCTT	2,00	0,37	Hs.91684	Homo sapiens cDNA FLJ20148 fis, clone COL08032, high
2674	GTTAAATCCTG	2,00	0,37	Hs.8881	ESTs, Weakly similar to RMS1_HUMAN REGULATOR OF MITO
2675	TAAGTAAAGTG	2,00	0,37	Hs.36529	ESTs
2676	ATCAAAAAAA	2,00	0,37	Hs.271699	polymerase (DNA directed) iota
2677	ACCTATTTGTG	2,00			v-maf musculoaponeurotic fibrosarcoma (avian) oncoge
	GAATGTTTTT	2,00	0,37	Hs.21432	SEX gene
2679	GTTGTGCCACT	2,00	0,37	Hs.196978	EST
2680	CACTTTAAAGT	2,00	0,37	Hs.194669	enhancer of zeste (Drosophila) homolog
2681	AAATAAAAAAT	2,00	0,37	Hs.151696	DKFZP727G051 protein
	AAGGTGAAGTA	2,00	0,37	Hs.146861	hypothetical protein FLJ20580
2683	CTTTAAGAAAG	2,00		Hs.13456	Homo sapiens clone 24747 mRNA sequence
2684	GTGGCTCTTGC	2,00	0,37	Hs.12243	hypothetical protein
	TCTCTACTAAA	16,00			PRO0365 protein
2686	CCCTTCCCCGG	6,00			Misshapen/NIK-related kinase
	ACCTATAAGTA	6,00			putative heme-binding protein

2689 CTCACTAGTGG         3,00         0,50 Hs.78683         ubiquitin specific protease 7 (herpes virus-associat           2690 GCGAAACCCCT         3,00         0,50 Hs.46468         chemokine (C-C motif) receptor 6           2691 GACTGTTGCTG         3,00         0,50 Hs.179902         Homo sapiens CTL1 gene           2692 CCACTGCAGTC         7,00         0,87 Hs.161554         hypothetical protein FLJ20159           2693 TCAACAAATTT         1,00         0,19 Hs.9315         HNDEL-iso protein           2694 GGACCTTAAT         1,00         0,19 Hs.89315         insulin receptor           2695 ATACTACACAGG         1,00         0,19 Hs.8867         cysteine-rich, angiogenic inducer, 61           2697 ATGCAACTGGG         1,00         0,19 Hs.8867         ESTs           2698 TTGCCTTCTCT         1,00         0,19 Hs.82906         CDC20 (cell division cycle 20, S. cerevisiae, homolo           2699 TTTTGTATGTC         1,00         0,19 Hs.79025         KIAA0096 protein           2701 TGGTAGGTTCT         1,00         0,19 Hs.76136         thioredoxin           2702 ACAAGATATTT         1,00         0,19 Hs.70202         hypothetical protein FLJ10897           2703 ACCAGCATCCT         1,00         0,19 Hs.67136         thioredoxin           2703 ACCAGCATCCTA         1,00         0,19 Hs.62711	2688	TTATACAAAAA	3,00	0.50	Hs.88558	FSTe
2690   CGAAACCCCT   3,00   0,50   Hs.46468   chemokine (C-C motif) receptor 6   2691   GACTGTTGCTG   3,00   0,50   Hs.179902   Home sapiens CTL1 gene   2692   CACCTGCAGTC   7,00   0,87   Hs.161554   hypothetical protein FLJ20159   2693   TCAACAAATTT   1,00   0,19   Hs.9315   HNOEL-iso protein   2694   GACCTTTAAT   1,00   0,19   Hs.90011   adenylosuccinate synthase   2695   GATACTTTGCA   1,00   0,19   Hs.8695   insulin receptor   2696   TATTGAACACG   1,00   0,19   Hs.8695   insulin receptor   2696   TATTGAACACG   1,00   0,19   Hs.8617   ESTs   2698   TTGCCTTCTCT   1,00   0,19   Hs.82906   CDC20 (cell division cycle 20, S. cerevisiae, homolo   2700   GAAATGCAGCC   1,00   0,19   Hs.78546   ATPase, Ca++ transporting, plasma   membrane 1   2701   TGGTAGGTTCT   1,00   0,19   Hs.78136   thioredoxin   2702   ACAAGATATTT   1,00   0,19   Hs.78136   thioredoxin   2702   ACAAGATATTT   1,00   0,19   Hs.78136   thioredoxin   2703   ACAAGATACCT   1,00   0,19   Hs.67816   ESTs   2703   ACAAGATACCT   1,00   0,19   Hs.6786   ESTs   2705   GTGCAGAGAGC   1,00   0,19   Hs.6786   ESTs   2705   GTGCAGAGAGC   1,00   0,19   Hs.6786   ESTs   2705   GTGCAGAGACC   1,00   0,19   Hs.6786   ESTs   2705   GTGCAGAGAGC   1,00   0,19   Hs.6786   ESTs   2707   CTATTTAGTTA   1,00   0,19   Hs.6781   ESTs   2708   CAAGTAGCAAA   1,00   0,19   Hs.86392   intersectin 1 (SH3 domain protein)   2708   CAAGTAGCAAA   1,00   0,19   Hs.87834   KIAA0281 gene product   2710   TGCACTGTGT   1,00   1,19   Hs.37834   KIAA0281 gene product   2710   TGCACTGTGGT   1,00   0,19   Hs.25585   ESTs   2711   CTATTCACTC   1,00   0,19   Hs.25685   ESTs   2712   TTTTTCCCC   1,00   0,19   Hs.25685   ESTs   2713   CTGAAACTGAC   1,00   0,19   Hs.25685   ESTs   2714   CTTACAAAAAA   1,00   0,19   Hs.25685   ESTs   2714   CTTACAACAAA   1,00   0,19   Hs.25685   ESTs   2715   GATTAAATAAT   1,00   0,19   Hs.25685   ESTs   2716   CTTTGTTAGAG   1,00   0,19   Hs.25685   ESTs   2717   TGAACCTGAC   1,00   0,19   Hs.22586   ESTs   2718   TAGCCAATGTT   1,00   0,19   Hs.22587   THOE IN TOTAG	2003	CIONOINGIGO	3,00	0,50	1 13.7 0000	
2691   CACTGTTGCTG   3,00   0,50   Hs.179902   Homo sapiens CTL1 gene   2692   CCACTGCACTC   7,00   0,87   Hs.161554   hypothetical protein FLJ20159   2693   TCAACAAAATTT   1,00   0,19   Hs.9315   HNOEL-iso protein   7,00   2694   GGACCTTTAAT   1,00   0,19   Hs.90011   adenylosuccinate synthase   2695   CATACTTTGCA   1,00   0,19   Hs.88695   insulin receptor   2696   TATTGAACACG   1,00   0,19   Hs.8867   Cysteine-rich, angiogenic inducer, 61   2697   ATGCAACTGGG   1,00   0,19   Hs.82906   CDC20 (cell division cycle 20, S. cerevisiae, homolo   CDC20   CATACTGGG   1,00   0,19   Hs.78546   ATPase, Ca++ transporting, plasma membrane 1   2701   TGGTAGGTTCT   1,00   0,19   Hs.78546   ATPase, Ca++ transporting, plasma membrane 1   2702   ACAAGATATTT   1,00   0,19   Hs.74122   caspase 4, apoptosis-related cysteine protease   2703   ACCAGCATCCT   1,00   0,19   Hs.6786   ESTs   CTGTCACGAT   1,00   0,19   Hs.6786   ESTs   2705   GTGCAGAGAGC   1,00   0,19   Hs.68392   intersectin 1 (SH3 domain protein)   2706   ATGTTACCTAA   1,00   0,19   Hs.62711   ESTs   2707   CTATTTAGTTA   1,00   0,19   Hs.62711   ESTs   2708   CAGAGAGAC   1,00   0,19   Hs.62711   ESTs   2708   CAGAGAGAC   1,00   0,19   Hs.8244   DKFZP434N061 protein   2712   TTTTGCCC   1,00   0,19   Hs.30524   ring finger protein 24   2711   CCATTTCACTC   1,00   0,19   Hs.23552   ring finger protein   24   2711   CCATTTCACTC   1,00   0,19   Hs.255298   ESTs   2715   GATTCACAAA   1,00   0,19   Hs.255298   ESTs   2715   GATTCACAAA   1,00   0,19   Hs.255298   ESTs   2717   TGAAGCCTGAC   1,00   0,19   Hs.255298   ESTs   2718   TGAGCCAAACT   1,00   0,19   Hs.25537   protein phosphatase 1, catalytic s	2690	GCGAAACCCCT	3.00	0.50	Hs 46468	
2693   TCAACAAATTT   1,00						
2693   TCAACAAATTT   1,00						
2694         GGACCTTTAAT         1,00         0,19 Hs.90011         adenylosuccinate synthase           2695         GATACTTTGAA         1,00         0,19 Hs.8867         insulin receptor           2696         TATTGAACAG         1,00         0,19 Hs.8617         ESTs           2698         TATGCAACTGGG         1,00         0,19 Hs.8617         ESTs           2698         TTGCCTTCTCT         1,00         0,19 Hs.79025         KIAA0096 protein           2700         GAAATGCAGCC         1,00         0,19 Hs.78546         ATPase, Ca++ transporting, plasma membrane           2701         TGGTAGGTTCT         1,00         0,19 Hs.76136         thioredoxin           2702         ACAAGATACTT         1,00         0,19 Hs.70202         hypothetical protein FLJ10897           2703         ACCAGCATCCT         1,00         0,19 Hs.6786         ESTs           2704         CCTGTCACGAT         1,00         0,19 Hs.66392         intersectin 1 (SH3 domain protein)           2705         GTGCAGAGAGC         1,00         0,19 Hs.592         ESTs           2706         CTATITAGTTA         1,00         0,19 Hs.2942         DKFZP434N061 protein           2706         TTATGAGAAA         1,00         0,19 Hs.2942         DKFZP434N061 protein						
2695   GATACTITIGCA						
2696   TATTGAACACG						
2697   ATGCAACTGGG						
2698   TTGCCTTCTCT						
Cerevisiae, homolo   Cerevis						
2699   TTTTGTATGTC	2090	TIGOCTICICI	1,00	0, 19	ns.62906	
2700   GAAATGCAGCC	2600	TTTCTATCTC	1.00	0.10	Ha 70025	
membrane 1     membrane 1       membrane 1						
2702   ACAAGATATTT   1,00						membrane 1
2703   ACCAGCATCCT   1,00   0,19   Hs.70202   hypothetical protein FLJ10897						
2704   CCTGTCACGAT   1,00   0,19   Hs.6786   ESTs     2705   GTGCAGAGAGC   1,00   0,19   Hs.66392   intersectin 1 (SH3 domain protein)     2706   ATGTTACCTAA   1,00   0,19   Hs.576   fucosidase, alpha-L- 1, tissue     2707   CTATTTAGTTA   1,00   0,19   Hs.576   fucosidase, alpha-L- 1, tissue     2708   CAAGTAGCAAA   1,00   0,19   Hs.4924   DKFZP434N061 protein     2709   TTAAGGGAAT   1,00   0,19   Hs.31463   KIAA0281 gene product     2710   TGCACTGTGGT   1,00   0,19   Hs.30524   ring finger protein 24     2711   CCATTCACTC   1,00   0,19   Hs.284467   EST     2712   TTTTCTCCCC   1,00   0,19   Hs.29583   CGI-81 protein     2713   CTGAACCTGAC   1,00   0,19   Hs.255372   hypothetical protein DKFZp564O1278     2714   TCTACAAAAAA   1,00   0,19   Hs.255298   ESTs     2715   GATTAAATAAT   1,00   0,19   Hs.256991   ESTs     2716   CTTGTTAGAA   1,00   0,19   Hs.246885   hypothetical protein FLJ20783     2718   TAGCCAATGTT   1,00   0,19   Hs.246885   hypothetical protein FLJ20783     2719   GCCTCCATAAA   1,00   0,19   Hs.24427   DKFZP566O1646 protein     2720   AATTGAGGTT   1,00   0,19   Hs.223395   myosin IXA     2720   AATTGAGGGT   1,00   0,19   Hs.223171   EST     2721   TTTAAAAAATTA   1,00   0,19   Hs.223171   EST     2722   AGCCATTGCAC   1,00   0,19   Hs.22144   Homo sapiens mRNA full length insert     2724   TATTCAGGGGC   1,00   0,19   Hs.21256   ESTs     2725   TATTCTGAACA   1,00   0,19   Hs.21256   ESTs     2726   CTTACAGCCAA   1,00   0,19   Hs.21256   ESTs     2727   AGTGACAAACT   1,00   0,19   Hs.195969   ESTs     2728   CTTACAGCCAA   1,00   0,19   Hs.195969   ESTs     2729   AGTGACAAACT   1,00   0,19   Hs.195969   ESTs     2721   Homo sapiens BAC clone RP11-505D1     2722   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2723   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2724   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2725   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2726   CTTAC	2702	ACAAGATATTT	1,00	0,19	Hs.74122	
2704   CCTGTCACGAT   1,00   0,19   Hs.6786   ESTs     2705   GTGCAGAGAGC   1,00   0,19   Hs.66392   intersectin 1 (SH3 domain protein)     2706   ATGTTACCTAA   1,00   0,19   Hs.576   fucosidase, alpha-L- 1, tissue     2707   CTATTTAGTTA   1,00   0,19   Hs.576   fucosidase, alpha-L- 1, tissue     2708   CAAGTAGCAAA   1,00   0,19   Hs.4924   DKFZP434N061 protein     2709   TTAAGGGAAT   1,00   0,19   Hs.31463   KIAA0281 gene product     2710   TGCACTGTGGT   1,00   0,19   Hs.30524   ring finger protein 24     2711   CCATTCACTC   1,00   0,19   Hs.284467   EST     2712   TTTTCTCCCC   1,00   0,19   Hs.29583   CGI-81 protein     2713   CTGAACCTGAC   1,00   0,19   Hs.255372   hypothetical protein DKFZp564O1278     2714   TCTACAAAAAA   1,00   0,19   Hs.255298   ESTs     2715   GATTAAATAAT   1,00   0,19   Hs.256991   ESTs     2716   CTTGTTAGAA   1,00   0,19   Hs.246885   hypothetical protein FLJ20783     2718   TAGCCAATGTT   1,00   0,19   Hs.246885   hypothetical protein FLJ20783     2719   GCCTCCATAAA   1,00   0,19   Hs.24427   DKFZP566O1646 protein     2720   AATTGAGGTT   1,00   0,19   Hs.223395   myosin IXA     2720   AATTGAGGGT   1,00   0,19   Hs.223171   EST     2721   TTTAAAAAATTA   1,00   0,19   Hs.223171   EST     2722   AGCCATTGCAC   1,00   0,19   Hs.22144   Homo sapiens mRNA full length insert     2724   TATTCAGGGGC   1,00   0,19   Hs.21256   ESTs     2725   TATTCTGAACA   1,00   0,19   Hs.21256   ESTs     2726   CTTACAGCCAA   1,00   0,19   Hs.21256   ESTs     2727   AGTGACAAACT   1,00   0,19   Hs.195969   ESTs     2728   CTTACAGCCAA   1,00   0,19   Hs.195969   ESTs     2729   AGTGACAAACT   1,00   0,19   Hs.195969   ESTs     2721   Homo sapiens BAC clone RP11-505D1     2722   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2723   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2724   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2725   AGTGACAAACT   1,00   0,19   Hs.18987   Homo sapiens BAC clone RP11-505D1     2726   CTTAC	2703	ACCAGCATCCT	1,00	0,19	Hs.70202	hypothetical protein FLJ10897
2706         ATGTTACCTAA         1,00         0,19         Hs.62711         ESTs           2707         CTATITAGTTA         1,00         0,19         Hs.576         fucosidase, alpha-L- 1, tissue           2708         CAAGTAGCAAA         1,00         0,19         Hs.4924         DKFZP434N061 protein           2709         TTTAAGGGAAT         1,00         0,19         Hs.30524         ring finger protein 24           2710         TGCACTGTGGT         1,00         0,19         Hs.284467         EST           2712         TTTTCCCCC         1,00         0,19         Hs.279583         CGI-81 protein           2712         TTTTCTCCCC         1,00         0,19         Hs.255372         hypothetical protein DKFZp564O1278           2714         TCTACAAAAAA         1,00         0,19         Hs.255298         ESTs           2715         GATTAAATAAT         1,00         0,19         Hs.255298         ESTs           2716         CTTTGTTAGAA         1,00         0,19         Hs.246885         hypothetical protein FLJ20783           2718         TAGCCAATGTT         1,00         0,19         Hs.24427         DKFZP566O1646 protein           2719         GCCTCCATAAA         1,00         0,19         Hs.22	2704	CCTGTCACGAT		0,19		
2706 ATGTTACCTAA         1,00         0,19 Hs.62711         ESTs           2707 CTATTTAGTTA         1,00         0,19 Hs.576         fucosidase, alpha-L- 1, tissue           2708 CAAGTAGCAAA         1,00         0,19 Hs.4924         DKFZP434N061 protein           2709 TTTAAGGGAAT         1,00         0,19 Hs.31463         KIAA0281 gene product           2710 TGCACTGTGGT         1,00         0,19 Hs.284467 EST           2711 CCATTTCACTC         1,00         0,19 Hs.279583 CGI-81 protein           2712 TTTTCTCCCC         1,00         0,19 Hs.279583 CGI-81 protein           2713 CTGAACCTGAC         1,00         0,19 Hs.255372 hypothetical protein DKFZp564O1278           2714 TCTACAAAAA         1,00         0,19 Hs.255298 ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.255485 ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.250691 ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.23395 myosin IXA           2721 TTTTAAAGAGG         1,00         0,19 Hs.221371 EST           2722 AGCCATTGCAC         1,00         0,19 Hs.22537 protein phosphatase 1, catalytic subunit beta isofo      <	2705	GTGCAGAGAGC				intersectin 1 (SH3 domain protein)
2707         CTATTTAGTTA         1,00         0,19         Hs.576         fucosidase, alpha-L- 1, tissue           2708         CAAGTAGCAAA         1,00         0,19         Hs.4924         DKFZP434N061 protein           2709         TTTAAGGGAAT         1,00         0,19         Hs.31463         KIAA0281 gene product           2710         TGCACTGTGGT         1,00         0,19         Hs.30524         ring finger protein 24           2711         CCATTTCACTC         1,00         0,19         Hs.284467         EST           2712         TTTTCTCCCC         1,00         0,19         Hs.279583         CGI-81 protein           2713         CTGAACCTGAC         1,00         0,19         Hs.255372         hypothetical protein DKFZp564O1278           2714         TCTACAAAAAA         1,00         0,19         Hs.255298         ESTs           2715         GATTAAATAAT         1,00         0,19         Hs.255298         ESTs           2717         TGAAGCCTTGC         1,00         0,19         Hs.246885         hypothetical protein FLJ20783           2718         TAGCCAATGTT         1,00         0,19         Hs.24427         DKFZP566O1646 protein           2719         GCCTCCATAAA         1,00         0,19 <td></td> <td></td> <td></td> <td>0,19</td> <td>Hs.62711</td> <td>ESTs</td>				0,19	Hs.62711	ESTs
2708 CAAGTAGCAAA         1,00         0,19 Hs.4924         DKFZP434N061 protein           2709 TTTAAGGGAAT         1,00         0,19 Hs.31463         KIAA0281 gene product           2710 TGCACTGTGGT         1,00         0,19 Hs.30524         ring finger protein 24           2711 CCATTTCACTC         1,00         0,19 Hs.284467 EST           2712 TTTTTCTCCCC         1,00         0,19 Hs.279583 CGI-81 protein           2713 CTGAACCTGAC         1,00         0,19 Hs.255372 hypothetical protein DKFZp564O1278           2714 TCTACAAAAAA         1,00         0,19 Hs.255298 ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.250691 ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.24427 DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883 ESTs           2721 TTTAAAGAGG         1,00         0,19 Hs.223171 EST           2723 TTTAAAAAATTA         1,00         0,19 Hs.223171 EST           2723 TTTAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA						
2709 TTTAAGGGAAT         1,00         0,19 Hs.31463         KIAA0281 gene product           2710 TGCACTGTGGT         1,00         0,19 Hs.30524         ring finger protein 24           2711 CCATTTCACTC         1,00         0,19 Hs.284467 EST           2712 TTTTTCTCCCC         1,00         0,19 Hs.279583 CGI-81 protein           2713 CTGAACCTGAC         1,00         0,19 Hs.255372 hypothetical protein DKFZp564O1278           2714 TCTACAAAAAA         1,00         0,19 Hs.255298 ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.25485 ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.250691 ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.24427 DKFZP566O1646 protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883 ESTs           2721 TTTAAAGAGG         1,00         0,19 Hs.22744 Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969						
2710 TGCACTGTGGT         1,00         0,19 Hs.30524         ring finger protein 24           2711 CCATTTCACTC         1,00         0,19 Hs.284467         EST           2712 TTTTCTCCCC         1,00         0,19 Hs.279583         CGI-81 protein           2713 CTGAACCTGAC         1,00         0,19 Hs.255372         hypothetical protein DKFZp564O1278           2714 TCTACAAAAAA         1,00         0,19 Hs.255298         ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.25691         ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.25691         ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.246885         hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.24427         DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395         myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22744         Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256         ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.2021         vesicle-associated membrane protein 1 (synaptobrevin						
2711 CCATTTCACTC         1,00         0,19 Hs.284467 EST           2712 TTTTCTCCCC         1,00         0,19 Hs.279583 CGI-81 protein           2713 CTGAACCTGAC         1,00         0,19 Hs.255372 hypothetical protein DKFZp564O1278           2714 TCTACAAAAAA         1,00         0,19 Hs.255298 ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.25485 ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.250691 ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.23395 myosin IXA           2719 GCCTCCATAAA         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22744 Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.2021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1						
2712         TTTTTCTCCCC         1,00         0,19         Hs.279583         CGI-81 protein           2713         CTGAACCTGAC         1,00         0,19         Hs.255372         hypothetical protein DKFZp564O1278           2714         TCTACAAAAAA         1,00         0,19         Hs.255298         ESTs           2715         GATTAAATAAT         1,00         0,19         Hs.250691         ESTs           2716         CTTTGTTAGAA         1,00         0,19         Hs.250691         ESTs           2717         TGAAGCCTTGC         1,00         0,19         Hs.246885         hypothetical protein FLJ20783           2718         TAGCCAATGTT         1,00         0,19         Hs.24427         DKFZP566O1646 protein           2719         GCCTCCATAAA         1,00         0,19         Hs.23395         myosin IXA           2720         AATTTGAGTGT         1,00         0,19         Hs.22883         ESTs           2721         TTTAAAAAATTA         1,00         0,19         Hs.223171         EST           2723         TTTAAAAATTA         1,00         0,19         Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724         TATTCTGAACA         1,00         0,19         Hs.2						
2713 CTGAACCTGAC         1,00         0,19 Hs.255372 hypothetical protein DKFZp564O1278           2714 TCTACAAAAAA         1,00         0,19 Hs.255298 ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.25485 ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.250691 ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.24427 DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883 ESTs           2721 TTTTAAAGAGG         1,00         0,19 Hs.223171 EST           2722 AGCCATTGCAC         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969 ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1						
2714 TCTACAAAAAA         1,00         0,19 Hs.255298 ESTs           2715 GATTAAATAAT         1,00         0,19 Hs.25485 ESTs           2716 CTTTGTTAGAA         1,00         0,19 Hs.250691 ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.24427 DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883 ESTs           2721 TTTTAAAGAGGG         1,00         0,19 Hs.22744 Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969 ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.195969 ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1	_					
2715         GATTAAATAAT         1,00         0,19         Hs.25485         ESTs           2716         CTTTGTTAGAA         1,00         0,19         Hs.250691         ESTs           2717         TGAAGCCTTGC         1,00         0,19         Hs.246885         hypothetical protein FLJ20783           2718         TAGCCAATGTT         1,00         0,19         Hs.24427         DKFZP566O1646 protein           2719         GCCTCCATAAA         1,00         0,19         Hs.23395         myosin IXA           2720         AATTTGAGTGT         1,00         0,19         Hs.22883         ESTs           2721         TTTAAAGAGGG         1,00         0,19         Hs.22744         Homo sapiens mRNA full length insert cDNA clone EURO           2722         AGCCATTGCAC         1,00         0,19         Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724         TATTCAGGGGC         1,00         0,19         Hs.21256         ESTs           2725         TATTCTGAACA         1,00         0,19         Hs.20021         vesicle-associated membrane protein 1 (synaptobrevin           2726         CTTACAGCCAA         1,00         0,19         Hs.195969         ESTs           2727         AGTGACAAACT						
2716 CTTTGTTAGAA         1,00         0,19 Hs.250691 ESTs           2717 TGAAGCCTTGC         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.24427 DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883 ESTs           2721 TTTAAAGAGG         1,00         0,19 Hs.22744 Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.223171 EST           2723 TTTAAAAAATTA         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969 ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1						
2717 TGAAGCCTTGC         1,00         0,19 Hs.246885 hypothetical protein FLJ20783           2718 TAGCCAATGTT         1,00         0,19 Hs.24427 DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395 myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883 ESTs           2721 TTTTAAAGAGG         1,00         0,19 Hs.22744 Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.223171 EST           2723 TTTAAAAATTA         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969 ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1						
2718 TAGCCAATGTT         1,00         0,19 Hs.24427         DKFZP566O1646 protein           2719 GCCTCCATAAA         1,00         0,19 Hs.23395         myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883         ESTs           2721 TTTTAAAGAGG         1,00         0,19 Hs.22744         Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.223171         EST           2723 TTTAAAAATTA         1,00         0,19 Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256         ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021         vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969         ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987         Homo sapiens BAC clone RP11-505D1						
2719 GCCTCCATAAA         1,00         0,19 Hs.23395         myosin IXA           2720 AATTTGAGTGT         1,00         0,19 Hs.22883         ESTs           2721 TTTTAAAGAGG         1,00         0,19 Hs.22744         Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.223171         EST           2723 TTTAAAAATTA         1,00         0,19 Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256         ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021         vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969         ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987         Homo sapiens BAC clone RP11-505D1						
2720 AATTTGAGTGT         1,00         0,19 Hs.22883         ESTs           2721 TTTTAAAGAGG         1,00         0,19 Hs.22744         Homo sapiens mRNA full length insert cDNA clone EURO           2722 AGCCATTGCAC         1,00         0,19 Hs.223171         EST           2723 TTTAAAAATTA         1,00         0,19 Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256         ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021         vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969         ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987         Homo sapiens BAC clone RP11-505D1						
2721         TTTTAAAGAGG         1,00         0,19         Hs.22744         Homo sapiens mRNA full length insert cDNA clone EURO           2722         AGCCATTGCAC         1,00         0,19         Hs.223171         EST           2723         TTTAAAAATTA         1,00         0,19         Hs.21537         protein phosphatase 1, catalytic subunit beta isofo           2724         TATTCAGGGGC         1,00         0,19         Hs.21256         ESTs           2725         TATTCTGAACA         1,00         0,19         Hs.20021         vesicle-associated membrane protein 1 (synaptobrevin           2726         CTTACAGCCAA         1,00         0,19         Hs.195969         ESTs           2727         AGTGACAAACT         1,00         0,19         Hs.18987         Homo sapiens BAC clone RP11-505D1						
2722 AGCCATTGCAC         1,00         0,19 Hs.223171 EST           2723 TTTAAAAATTA         1,00         0,19 Hs.21537 protein phosphatase 1, catalytic subunit beta isofo           2724 TATTCAGGGGC         1,00         0,19 Hs.21256 ESTs           2725 TATTCTGAACA         1,00         0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin           2726 CTTACAGCCAA         1,00         0,19 Hs.195969 ESTs           2727 AGTGACAAACT         1,00         0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1						Homo sapiens mRNA full length insert
2723TTTAAAAATTA1,000,19Hs.21537protein phosphatase 1, catalytic subunit beta isofo2724TATTCAGGGGC1,000,19Hs.21256ESTs2725TATTCTGAACA1,000,19Hs.20021vesicle-associated membrane protein 1 (synaptobrevin2726CTTACAGCCAA1,000,19Hs.195969ESTs2727AGTGACAAACT1,000,19Hs.18987Homo sapiens BAC clone RP11-505D1	2722	AGCCATTGCAC	1 00	0.19	Hs 223171	
beta isofo   2724 TATTCAGGGGC						
2725 TATTCTGAACA 1,00 0,19 Hs.20021 vesicle-associated membrane protein 1 (synaptobrevin 2726 CTTACAGCCAA 1,00 0,19 Hs.195969 ESTs 2727 AGTGACAAACT 1,00 0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1						beta isofo
(synaptobrevin   2726 CTTACAGCCAA   1,00   0,19 Hs.195969 ESTs   2727 AGTGACAACT   1,00   0,19 Hs.18987   Homo sapiens BAC clone RP11-505D1						
2727 AGTGACAAACT 1,00 0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1	2725	TATTCTGAACA	1,00			(synaptobrevin
2727 AGTGACAAACT 1,00 0,19 Hs.18987 Homo sapiens BAC clone RP11-505D1	2726	CTTACAGCCAA	1,00	0,19		
	2727	AGTGACAAACT				Homo sapiens BAC clone RP11-505D17 from 7p22-p21
2728 TTGGCAAAAA 1,00 0,19 Hs.18136 inicotinamide nucleotide	2728	TTGGCAAAAA	1.00	0.19	Hs.18136	

	<u> </u>	1		<del></del>	lt-on-bud-oggnoo
2720	CCTCACACTCA	4.00	0.40	Up 474400	transhydrogenase
	GCTGACACTGA	1,00			chloride channel 3
	AATCAGAATTT	1,00			KIAA0941 protein
	TGACCATTTCT	1,00		Hs.171952	
2732	CCAAAGCTTCC	1,00	0,19	Hs.171637	ESTs, Weakly similar to KIAA1317
		1 2 2			protein [H.sapiens]
2733	ATGTTCAATTT	1,00	0,19	Hs.1710	ATP-binding cassette, sub-family B
		4 00		11 404554	(MDR/TAP), member
-	CCACACAAAA	1,00			hypothetical protein FLJ20159
	GCCGGGCGCAG	1,00		Hs.160821	
2736	TCATCACATTA	1,00	0,19	Hs.155566	CASP2 and RIPK1 domain containing
					adaptor with death
	TTACCCAGGCT	1,00			hypothetical protein FLJ11193
2738	CCTCAGTTTGG	1,00	0,19	Hs.140944	DNA segment on chromosome X
					(unique) 2654 expressed
	CACCTATAGTC	1,00			Rho GTPase activating protein 1
	GGTGCTTCTTT	1,00		Hs.129692	
	GAGTCCATAAA	1,00		Hs.126266	
	CTAACAGGATT	1,00			choline/ethanolaminephosphotransferase
	TCGAAACCCCA	1,00	0,19	Hs.117582	CGI-43 protein
2744	AAGGAGTGAAA	1,00	0,19	Hs.111092	ESTs, Weakly similar to gonadotropin
					inducible trans
	ATCCCTGTGAG	1,00		Hs.105820	
2746	CTCCATCGGCT	5,00	0,70	Hs.65238	95 kDa retinoblastoma protein binding
					protein; KIAA0
	GTGGAACCCCG	5,00		Hs.270796	
	GACCACGAATA	4,00			cathepsin H
$\overline{}$	CCCTTGTGACT	4,00			cytochrome c oxidase subunit VIc
2750	GACCGCGGCTT	4,00	0,60	Hs.110903	claudin 5 (transmembrane protein
					deleted in velocard
2751	CTACCAGGCCT	6,00	0,78	Hs.54457	CD81 antigen (target of antiproliferative
					antibody 1
	TTCCGGTTCCA	18,00			nucleobindin 1
2753	GATAGCACAGT	27,00	2,10	Hs.103391	Human insulin-like growth factor binding
					protein 5 (
	TACTGTATGTC	3,00	0,49	Hs.5867	KIAA0851 protein
2755	TAATTTCTCAA	3,00	0,49	Hs.5306	Homo sapiens mRNA; cDNA
					DKFZp586F1122 (from clone DK
2756	GCAAAACCCCC	3,00	0,49	Hs.277209	EST, Weakly similar to ALU7_HUMAN
					ALU SUBFAMILY SQ S
	GTGCGGTACCT	3,00			microspherule protein 1
	TACTGCAAAAA	3,00		Hs.24557	DKFZP434H018 protein
2759	TGCCTTAGTAA	3,00	0,49	Hs.13015	ESTs, Highly similar to MTJ1_MOUSE
					DNAJ PROTEIN HOMO
	TACTGTGATGT	3,00			chromosome 11 open reading frame 15
	GGGGCTGGAGG	5,00			KIAA0620 protein
	AACGCTGCGAA	4,00			KIAA0876 protein
	ATGGTGGGCAC	4,00			zinc finger protein 36 (KOX 18)
2764	TTGACTITTGC	2,00	0.36	Hs.61441	KIAA1311 protein

0305	COTOTALTOGO	0.001		11- 07075	It was the stirred markets and 100 440
	CCTGTAATCCG	2,00			hypothetical protein FLJ20413
	CTGTGAAAAA	2,00			kallikrein 2, prostatic
	CAGTGCAGTAT	2,00			Ank, mouse, homolog of
	AGCCATTGTGC	2,00		Hs.163860	
	GGCAGGGTCGA	2,00			Homo sapiens cDNA FLJ10158 fis, clone HEMBA1003463
	TAATTTTGAAA	12,00			CGI-43 protein
	TCTGGCATAGC	6,00			ESTs, Weakly similar to 159365 ubiquitin conjugating
	TTTTATGGAAT	12,00	_	Hs.77269	guanine nucleotide binding protein (G protein), alph
	GATTTTTAAAA	3,00			ralA binding protein 1
	GACGGCCAGAG	3,00		Hs.74649	cytochrome c oxidase subunit VIc
	TTCTCTGCTCA	3,00		Hs.21907	histone acetyltransferase
	TGGAAGGACCG	7,00		Hs.5086	Human DNA from overlapping chromosome 19 cosmids R31
	TGGGACTCCAG	4,00			Homo sapiens mRNA; cDNA DKFZp586E2023 (from clone DK
	TGGAATTCCCT	4,00			phosphatidylinositol 4-kinase, catalytic, beta polyp
	GCTCTCCCCCC	6,00			nuclear receptor subfamily 1, group H, member 2
2780	AGTGGAGGGAA	5,00	0,68	Hs.43509	ataxin 2 related protein
2781	AGCCACCGCAC	13,00	1,23	Hs.42612	ESTs
2782	TCTGAAGTCAA	8,00	0,90		inhibitor of DNA binding 2, dominant negative helix-
	AACGCGAACAC	7,00			squamous cell carcinoma antigen recognised by T cell
	GTGGCGCACAC	15,00	_		ESTs, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ
	ATGGTGCCACT	8,00	0,90	Hs.107479	KIAA0738 gene product
	TCAGAAAAAA	2,00			KIAA0650 protein
2787	TCCCCGTGGCT	2,00			KIAA0018 gene product
	AATAGGGGAAA	2,00			KIAA1075 protein
	AGTCCTTATGC	2,00			Homo sapiens cDNA FLJ10092 fis, clone HEMBA1002349
	TCCTTGGACCT	2,00	0,35	Hs.274550	proline oxidase homolog
2791	ATTITGCTTGG	2,00	0,35	Hs.249495	heterogeneous nuclear ribonucleoprotein A1
2792	CTTGTGTTATA	2,00	0,35	Hs.155024	B-cell CLL/lymphoma 6 (zinc finger protein 51)
2793	CAGCTTAATTA	2,00	0,35	Hs.13996	ESTs
	TTGGTCAGGTT	2,00		Hs.113111	Homo sapiens familial Mediterranean fever locus regi
2795	GTAATTCTCAA	2,00	0,35	Hs.107001	ESTs, Highly similar to 45kDa splicing factor [H.sap
2796	GCCCACAGCC	5,00	0,67		putative nuclear protein
	GTATACAACAA	3,00			steroid dehydrogenase homolog
	AGCTAGGGAAG	3,00			KIAA0440 protein

2700	TAATTGAAATA	3,00	0.48	He 146669	KIAA1253 protein
	TGATGTGATCA	3,00		Hs.12272	beclin 1 (coiled-coil, myosin-like BCL2-
2000	I GAIGIGAICA	3,00	0,40	ПS. 12212	interacting
2801	TGTGAGCCCCT	9,00	0.06	He 102048	enigma (LIM domain protein)
	TACAAGTTTTG	1,00			ESTs
	GGTACCAAAAA	1,00		Hs.9450	zinc finger protein 84 (HPF2)
	CCTGGCTCAAA	1,00		Hs.93127	Homo sapiens mRNA; cDNA
2004	CCTGGCTCAAA	1,00	0, 10	115.93 121	DKFZp586l0523 (from clone DK
2805	TAAGCCCTTTT	1,00	0.18	Hs.90606	15 kDa selenoprotein
	AGACCAATGAA	1,00		Hs.9029	DKFZP434G032 protein
	TGTTTGACTGT	1,00		Hs.8834	ring finger protein 3
	GACCAGCTGCC	1,00		Hs.84728	Kruppel-like factor 5 (intestinal)
	AGTTGAAGAAG	1,00		Hs.82120	nuclear receptor subfamily 4, group A,
					member 2
	AGTAGAAGAAT	1,00		Hs.74561	alpha-2-macroglobulin
2811	GAGCCCAGCCC	1,00	0,18	Hs.72885	azurocidin 1 (cationic antimicrobial
^					protein 37)
2812	CACATTTAATT	1,00	0,18	Hs.67619	Homo sapiens cDNA FLJ10533 fis, clone NT2RP2001056
2813	ACATATTGAGC	1,00	0,18	Hs.41086	hypothetical protein FLJ20067
2814	TTTGACTGATA	1,00	0,18	Hs.41086	hypothetical protein FLJ20067
2815	ACCAATACACT	1,00		Hs.29488	ESTs
2816	GTGAGACCCTA	1,00	0,18	Hs.280237	EST
2817	AAGATATTCTC	1,00	0,18	Hs.279934	Homo sapiens mRNA; cDNA
_					DKFZp434I0835 (from clone DK
2818	GTGGTGCACCC	1,00	0,18	Hs.278709	EST, Weakly similar to A46010 X-linked
					retinopathy p
2819	TTCTTGCAGCA	1,00	0,18	Hs.273234	Human DNA sequence from clone
					223H9 on chromosome 22
	CTTAATATATT	1,00			tumor differentially expressed 1
	GTGATACCCCA	1,00		Hs.270302	
	GAAAATATTCC	1,00			KIAA1099 protein
	ATGAAACCTTG	1,00		Hs.253488	
	CAAGATGAACC	1,00			E1A binding protein p300
2825	ACAAGAAAAA	1,00	0,18		microtubule-associated protein, RP/EB
					family, member
	TTTCGATTTTT	1,00	0,18	Hs.225951	topoisomerase-related function protein 4
2827	ATAAAACCCTG	1,00	0,18	Hs.223317	EST, Weakly similar to ALUC_HUMAN !!!! ALU CLASS C W
2828	GTGGCACGCAT	1,00	0,18	Hs.207823	Human EST clone 53125 mariner
					transposon Hsmar1 sequ
	AGGTGTTTCTT	1,00	0,18	Hs.19978	arylsulfatase D
	CTTTCTGTTAT	1,00			plasma glutamate carboxypeptidase
2831	CTCCAAATCCC	1,00	0,18	Hs.197289	rab3 GTPase-activating protein, non-
					catalytic subuni
2832	GGGGGCTCCTT	1,00	0,18	Hs.19718	protein tyrosine phosphatase, receptor
					type, U
	CTTGAAAAAAA	1,00			CGI-76 protein
2834	AAGGTGAGGAC	1,00	0,18	Hs.184216	DKFZP564C152 protein

2835	CAGGGGGTTCA	1,00	0.19	He 193630	hypothetical protein FLJ10210
-					
	CCATTATTCTT	1,00			RNA binding motif protein 3
	теттттете	1,00			Homo sapiens cDNA FLJ11066 fis, clone PLACE1004885
2838	TGATCCATCCT	1,00	0,18	Hs.178121	KIAA0626 gene product
2839	ATTAGTGTTGG	1,00	0,18	Hs.177781	superoxide dismutase 2, mitochondrial
2840	GCTATATTTCA	1,00	0,18	Hs.173571	KIAA1053 protein
2841	GTGCTATCCTG	1,00	0,18	Hs.172330	ESTs, Weakly similar to Wiskott-Aldrich Syndrome pro
2842	GGGAGACCCTG	1,00	0,18	Hs.169401	apolipoprotein E
2843	CTTTCAATGTT	1,00			vascular Rab-GAP/TBC-containing
	CCATTCTTTTC	1,00		Hs.162685	ESTs, Moderately similar to similar to KIAA0855 [H.s
2845	CCTCTTTAAAG	1,00	0,18	Hs.156016	KIAA0140 gene product
2846	AAGAAGGGATC	1,00			ESTs, Weakly similar to CNBP_HUMAN CELLULAR NUCLEIC
2847	AACCGCACCCA	1,00	0,18	Hs.142003	ESTs, Weakly similar to The KIAA0149 gene product is
2848	GTAAATGAGTA	1,00	0,18	Hs.138717	ESTs
2849	TAAACAAGCAA	1,00	0,18	Hs.135917	hypothetical protein DKFZp761D1823
2850	TGTATGTGGTA	1,00	0,18	Hs.13299	Homo sapiens mRNA; cDNA DKFZp761M0111 (from clone DK
2851	AGCCAGGGTAA	1,00	0.18	Hs.131525	KIAA0998 protein
	TTTTGATCACT	1,00			KIAA0560 gene product
Ī	CAGCTGGCACT	1,00		Hs.124418	
	TTAAGGGATGA	1,00		Hs.11177	
_	CAGAATGAGCC	1,00			KIAA0712 gene product
	TGACCACCTAC	1,00			DKFZP547E2110 protein
	GTGGCAGCCGG	1,00			KIAA0821 protein
Ì	GAAGCCATTGT	1,00			ATPase, Ca++ transporting, type 2C, member 1
2859	TATATGGATGG	1,00	0,18	Hs.106309	Friend of GATA2
2860	GTGCCCGGCAC	1,00			ESTs, Weakly similar to zinc finger protein zfp6 [H.
	GACCTTATTTT	1,00	0,18	Hs.100217	chromosome 17 open reading frame 1B
2862	CTGGGCCAGCC	4,00			vesicle-associated membrane protein 5 (myobrevin)
2863	GCCAAGGGGCC	7,00			oxoglutarate dehydrogenase (lipoamide)
2864	TCCTACAATCT	3,00			f-box and leucine-rich repeat protein 11
	TAACTTAAGCA	3,00			CGI-127 protein
	TGTAAAAAAAA	3,00	0,47	Hs.112743	synaptonemal complex protein 1
2867	TACCCTAGAAC	27,00	1,88	Hs.252588	Homo sapiens mRNA; cDNA DKFZp564F172 (from clone DKF
2868	TCAGCAATAAA	5,00	·		Homo sapiens cDNA FLJ20801 fis, clone ADSU01303
2869	CCTGTAATCCA	10,00	0,99	Hs.253369	
2870	TTGTGGAAATC	2,00		Hs.7807	ESTs, Weakly similar to predicted using Genefinder [
2871	ATGGGCACTGA	2,00	0.35	Hs.274201	60S acidic ribosomal protein PO

2872	GCTGACGGAAA	2,00	0,35	Hs.226377	phosphate cytidylyltransferase 2,
					ethanolamine
2873	GTGCCTCGGAG	2,00			Homo sapiens cDNA FLJ20418 fis, clone KAT02427
2874	ATGCTAAAAAA	5,00	0,65	Hs.116455	
2875	CTGCTGCCCCA	3,00	0,46	Hs.82407	Homo sapiens HSPC296 mRNA, partial cds
2876	TGAGTGGTAGC	3,00	0,46	Hs.77886	lamin A/C
2877	GTGAGACCTTG	3,00		Hs.246469	EST
2878	GTAGCAGGTGT	11,00	1,04	Hs.140452	cargo selection protein (mannose 6 phosphate recepto
2879	GTCTGGGGGAT	6,00	0,72	Hs.6721	lysophospholipase-like
2880	TTCCACTAACC	13,00	1,15	Hs.79706	plectin 1, intermediate filament binding protein, 50
2881	TGAAGCAGTAA	8,00	0,85	Hs.100407	Homo sapiens mRNA; cDNA DKFZp564H2416 (from clone DK
2882	ACCGCTTGTTT	4,00			inositol 1,4,5-triphosphate receptor, type 3
2883	CTGGCCAGGCT	4,00	0,56	Hs.182611	solute carrier family 11 (proton-coupled divalent me
2884	AGATTCAAACT	11,00	1,03	Hs.14368	SH3 domain binding glutamic acid-rich protein like
2885	AGTTGTCCCGG	6,00	0,71	Hs.239686	KIAA0462 protein
2886	CTGACTGTCCT	1,00	0,17	Hs.84298	CD74 antigen (invariant polypeptide of major histoco
2887	ACTGTATTGGA	1,00	0,17	Hs.83114	crystallin, zeta (quinone reductase)
2888	CAAATAACAAG	1,00	0,17	Hs.8262	lysosomal-associated membrane protein 2
2889	ACTTGTAATTA	1,00	0,17	Hs.80624	ESTs
2890	GACGACATTCG	1,00	0,17	Hs.79709	phosphotidylinositol transfer protein
2891	TACAAAGATGT	1,00	0,17		ESTs
2892	TTTTCCCCTGA	1,00	0,17	Hs.7822	Homo sapiens mRNA; cDNA DKFZp564C1216 (from clone DK
2893	CTTGGATGGCT	1,00	0,17	Hs.7314	KIAA0614 protein
2894	GTAAAATGCAA	1,00	0,17	Hs.66744	twist (Drosophila) homolog (acrocephalosyndactyly 3;
2895	CGATCACTGAC	1,00	0,17	Hs.6434	hypothetical protein DKFZp761F2014
2896	ATTGAGTCCAG	1,00			ESTs
	GGGCTTACTGT	1,00	0,17	Hs.5472	hypothetical protein FLJ20173
	TACCTATTGTG	1,00	0,17	Hs.48958	ESTs
2899	CTTTAAGGTTT	1,00	0,17	Hs.48827	Homo sapiens mRNA; cDNA
					DKFZp761D0223 (from clone DK
	ATGCTCTATTT	1,00			KIAA1286 protein
	GCAGGGAAATA	1,00		Hs.29263	ESTs
	AAAATGACAGA	1,00			signal sequence receptor, gamma (translocon-associat
	AGCCCACTGCT	1,00			KIAA0544 protein
2904	GGCCAGTGAGG	1,00	0,17	Hs.279899	tumor necrosis factor receptor superfamily, member 1

וסחתבו	CCTTGCCTGAA	1.00	0.17	Un 270424	SCO (cytochrome oxidase deficient,
2905	CCTTGCCTGAA	1,00	0,17	ITS.210431	yeast) homolog 2
2006	AAAGTGTATTT	1.00	0.17	He 274206	KIAA1423 protein
	CAGCTTTGCTG	1,00 1,00		Hs.25248	
		1,00		Hs.25005	
-	TGAGCACTCGA			Hs.249622	
	CCTGTAATTTC	1,00			
	AGTCACCAGCA	1,00			CREB binding protein (Rubinstein-Taybi syndrome)
	GGTACACATAC	1,00		Hs.225568	
	TGAAAGTGATT	1,00			KIAA0294 gene product
2913	CAGAGGCGTCC	1,00			eukaryotic translation initiation factor 2C, 2
2914	TTTGTTGTTGA	1,00	0,17		protein kinase, interferon-inducible double stranded
2915	TGCCCAACTTC	1,00	0.17		KIAA1319 protein
	TAGTATGGCTA	1,00			hypothetical protein
	CTATGTCTTTA	1,00			SWI/SNF related, matrix associated,
	,	.,	-,		actin dependent
2918	CAAGTAACTAG	1,00	0.17	Hs.172199	adenylate cyclase 7
	CTTCCTTGTAG	1,00			hypothetical protein FLJ11085
	GAAGATTGAGA	1,00			signal transducer and activator of
		.,	-,		transcription 5A
2921	CCGTGAAGTTT	1,00	0.17	Hs.159448	
	GTGATGTGCGC	1,00		Hs.158455	
	TGATTACAATC	1,00			voltage-dependent anion channel 1
	TTTCAATACCA	1,00		Hs.127006	
	GCAACCGTACT	1,00		Hs.126388	
_	CTATGTTCTGT	1,00		Hs.11494	
	CCTAACGATAC	1,00		Hs.11317	
	TAAAGGCTTTT	1,00			hypothetical protein FLJ10498
	GACACAGTTAT	1,00			ALR-like protein
	TGTAGAAAAA	10,00			tubulin, beta polypeptide
	CTCATATGTTA	3,00			yes-associated protein 65 kDa
	GGAGCCAGGCC	3,00			glutathione S-transferase theta 1
	GTGAAACACCA	3,00		Hs.231777	
	GCTCTCGGCGG	3,00		Hs.183994	protein phosphatase 1, catalytic subunit, alpha isof
2935	CAGATTGCTGA	5,00	0,63	Hs.21537	protein phosphatase 1, catalytic subunit, beta isofo
2936	TCCACTACCAA	4,00	0.55	Hs.18885	CGI-116 protein
	TATCCTGGCTC	4,00			erythrocyte membrane protein band 7.2
					(stomatin)
	CCCAGCCTGAA	2,00			ESTs
2939	TCTTTTAAAA	2,00	0,34	Hs.748	fibroblast growth factor receptor 1 (fms- related tyr
2940	AGGAGATGGAG	2,00	0,34	Hs.73987	CDC-like kinase 3
_	CTGCAGAATAT	2,00		Hs.4310	eukaryotic translation initiation factor 1A
	ATTITCCTTAG	2,00			APR-1 protein
	TATTTATTCAA	2,00			KIAA1228 protein

2011	CCGATTTTTAC	2,00	0.34	He 26570	hypothetical protein FLJ20422
	AGCCTCGGCCA	2,00			Rho guanine nucleotide exchange factor
2945	AGCCTCGGCCA	2,00	0,34	NS.20220U	(GEF) 1
2946	GGACCACCCAA	2,00	0,34	Hs.239298	microtubule-associated protein 4
2947	CGAGTGAGAGG	2,00	0,34	Hs.206259	Homo sapiens mRNA for KIAA1190
					protein, partial cds
2948	GTGAACCCTGT	2,00		Hs.204169	
2949	TGTGCATCTTG	2,00	0,34	Hs.18878	ESTs, Weakly similar to dJ876B10.4
					[H.sapiens]
2950	TGTAGTATTTG	2,00	0,34	Hs.18842	protein kinase C and casein kinase
					substrate in neur
-	TCTATCTCAGG	2,00			endothelin converting enzyme 1
2952	CGGGGTGGCCG	2,00	0,34	Hs.1584	cartilage oligomeric matrix protein
					(pseudoachondrop
-	CAGTGTATATA	2,00			HSPC040 protein
	ACTITAGATGG	8,00			collagen, type VI, alpha 3
2955	GCCCAAGGACC	31,00	1,90		filamin A, alpha (actin-binding protein-
		100.00			280)
	TGCACGTTTTC	189,00			ribosomal protein L32
	CATTATAACTT	5,00			hypothetical protein
	GACCCTAGCTC	5,00			polyglutamine binding protein 1
-	AGCCACTGCAC	23,00		Hs.122126	
2960	AGGGGCCGGGG	4,00	0,55		aldehyde dehydrogenase 4 (glutamate
0004	40044004	4 00	0.55	11- 470004	gamma-semialdehy
2961	AGGAAGGAACA	4,00	0,55	HS.1/3064	v-erb-b2 avian erythroblastic leukemia
2062	COTAAACTCAA	3.00	0.45		viral oncogen
2902	CCTAAACTCAA	3,00	0,45		uncharacterized hematopoietic
2062	теттсссттте	3,00	0.45		stem/progenitor cells MAX-interacting protein 1
	GTTCATAGTAT	3,00			N-terminal acetyltransferase complex
2904	GITCATAGIAI	3,00	0,45	ПS. 109200	ard1subunit
	AACTAACAAAA	59,00			ribosomal protein S27a
2966	AAGGTGCCTCC	6,00	0,69		heat shock 70kD protein 5 (glucose-
·					regulated protein
	GGGCTTCTGT	5,00			protease, cysteine, 1 (legumain)
	GTGGTGGACAC	13,00			hypothetical protein PRO1847
2969	CCGTGACTCTG	19,00	1,35	Hs.155712	Homo sapiens mRNA; cDNA
	70010155				DKFZp586O2223 (from clone DK
2970	TGCAGATGGTT	6,00	0,69	Hs.3628	mitogen-activated protein kinase kinase
007/	AAA00A4#046	0.05		11 70070	kinase kinas
	AAAGGAATGAG	2,00			KIAA0652 gene product
29/2	ATAATAAAGCT	2,00	0,33	Hs.37682	retinoic acid receptor responder
2072	CTCCACAAATT	200	0.00	LI- OCCOOR	(tazarotene induced
29/3	CTGCAGAAATT	2,00	0,33	TIS. 2002U5	Homo sapiens clone 23938 mRNA
2074	ATCCATCCCCT	2.00	0.22	Un 205244	sequence
	ATCGATCGCCT	2,00			KIAA1067 protein
29/5	CCAGTAACCCC	2,00	0,33	ms.230442	ESTs, Moderately similar to
2076	CACTITICTGT	2,00	0.33	Hs.23786	ALU7_HUMAN ALU SUBFAMILY ESTs
2010	UNUTTICIDI	2,00	0,33	113.23700	LU19

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2977	GTGGCATACAC	2,00	0,33	Hs.223603	Homo sapiens mRNA; cDNA DKFZp761D09121 (from clone D
2978	GTCCCAGGATT	2,00	0,33	Hs.158084	peroxisome receptor 1
2979	ATGTCTTCGTT	2,00	0,33	Hs.144926	ESTs
2980	AAAAGGAGATC	2,00	0,33	Hs.144904	nuclear receptor co-repressor 1
2981	GCGACCAACAT	5,00		Hs.4055	chromosome 21 open reading frame 50
2982	ATCCGCCTGCT	5,00	0,61	Hs.279904	ESTs
2983	GTGAAACCCAT	5,00	0,61	Hs.17311	hypothetical protein FLJ20004
2984	GACCACCTTTA	3,00	0,44	Hs.83551	microfibrillar-associated protein 2
2985	GAGCTTACATT	3,00			ESTs, Weakly similar to ALUC_HUMAN IIII ALU CLASS C
2986	GTGAGACCTCA	3,00			novel SH2-containing protein 1
2987	GAAACAAAATG	3,00	0,44	Hs.14896	DHHC1 protein
2988	GGTGAAGACAA	10,00	0,91	Hs.26951	KIAA0375 gene product
2989	TTGCTGTGTGA	4,00	0,53		actin binding protein; macrophin (microfilament and
	GTCTAGAATCT	5,00			vitamin A responsive; cytoskeleton related
2991	GCGAAACCCTG	62,00			v-erb-b2 avian erythroblastic leukemia viral oncogen
2992	CTTCCTGTACA	7,00	0,74	Hs.273237	postmeiotic segregation increased 2-like 8
2993	GCTATGGTTTC	1,00	0,17		ESTs, Weakly similar to unnamed protein product [H.s
2994	GTGCAGTTAGC	1,00			CGI-56 protein
2995	GTGCTATTATT	1,00	0,17		myeloid cell leukemia sequence 1 (BCL2-related)
	GACCTTAAGGC	1,00	0,17	Hs.8562	hypothetical protein FLJ20374
2997	AGCAGTCCAAC	1,00			acidic protein rich in leucines
2998	CCACTCCAAAA	1,00	0,17		defender against cell death 1
2999	TTGTACAACTG	1,00			baculoviral IAP repeat-containing 1
3000	TCTTTTTCAG	1,00	0,17	Hs.75209	protein kinase (cAMP-dependent, catalytic) inhibitor
3001	ACGCCGCCCAA	1,00	0,17	Hs.7306	secreted frizzled-related protein 1
	TGGTTTAAAAA	1,00		Hs.6298	KIAA1151 protein
	TAATTAAAAAA	1,00		Hs.60293	ESTs
3004	CTTCATAACCA	1,00	0,17	Hs.55896	ESTs
	AGAAGCAAGAG	1,00			IL-17B receptor
	TGTGGCCTCCA	1,00	0,17	Hs.5258	chromosome 11 open reading frame2
3007	GGCAGAAGATA	1,00		Hs.49349	beta-site APP-cleaving enzyme
3008	AGATAATCTGT	1,00	0,17	Hs.46571	SH3 domain protein 1B
	CATTTTGGATG	1,00		Hs.3994	ESTs
3010	GTGAAGAGTAA	1,00	0,17	Hs.38750	ESTs, Weakly similar to mSin3A associated polypeptid
3011	ATCAGTATGTG	1,00	0,17	Hs.278554	heterochromatin-like protein 1
3012	AGAATCCCTTG	1,00		Hs.277073	
3013	GAAGAAAAAA	1,00	0,17	Hs.274429	Homo sapiens cDNA FLJ11192 fis, clone PLACE1007618,
3014	ACTGATCTTGA	1,00	0,17	Hs.27268	Homo sapiens mRNA; cDNA

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DIKFZp564N198 (from clone DKF	3015   ACCCTTTTTT   1,00   0,17   Hs.257486   hypothetical protein FLJ2006   3016   TTGTAAAAAA   1,00   0,17   Hs.251377   taxol resistance associated g   3017   GGGGAAAGCAG   1,00   0,17   Hs.242262   ESTs   3018   TAACTGTCAAA   1,00   0,17   Hs.238205   hypothetical protein PRO201   3019   GTGGCTGATGC   1,00   0,17   Hs.238118   EST   3020   AAATTTGTATG   1,00   0,17   Hs.235445   Homo sapiens mRNA; cDNA   DKFZp761G2312 (from clone   3021   AGCCACCAGGC   1,00   0,17   Hs.230396   EST   3022   AAGGACATTCG   1,00   0,17   Hs.214982   laminin, gamma 1 (formerly L   3023   CCATAGATTTC   1,00   0,17   Hs.194035   KIAA0737   gene product   3024   TTCTCAATACA   1,00   0,17   Hs.193164   ESTs   3025   GCTTGTATGAG   1,00   0,17   Hs.18387   transcription factor AP-2   alphenancer   3027   GGAATTTGCTT   1,00   0,17   Hs.18387   transcription factor AP-2   alphenancer   3029   GTGACGCCCGG   1,00   0,17   Hs.183745   ESTs   3029   GTGACGCCCCG   1,00   0,17   Hs.173661   tubulin, beta polypeptide   3030   CAATGAGCATC   1,00   0,17   Hs.173993   RNA binding motif protein 6   3031   AATGTACCTGG   1,00   0,17   Hs.173422   cAMP responsive element binprotein 3 (luman)   3032   GCCCTCATTAA   1,00   0,17   Hs.169370   FYN oncogene related to SR	
3016   TGTAAAAAAA	3016   TTGTAAAAAAA	
3018   TAACTGTCAAA	3017 GGGGAAAGCAG         1,00         0,17 Hs.242262 ESTs           3018 TAACTGTCAAA         1,00         0,17 Hs.238205 hypothetical protein PRO201           3019 GTGGCTGATGC         1,00         0,17 Hs.238118 EST           3020 AAATTTGTATG         1,00         0,17 Hs.235445 Homo sapiens mRNA; cDNA DKFZp761G2312 (from clone DKFZp761G2312 (from clone DKFZp761G2312)           3021 AGCCACCAGGC         1,00         0,17 Hs.230396 EST           3022 AAGGACATTCG         1,00         0,17 Hs.214982 laminin, gamma 1 (formerly L 2022)           3023 CCATAGATTTC         1,00         0,17 Hs.194035 KIAA0737 gene product           3024 TTCTCAATACA         1,00         0,17 Hs.193164 ESTs           3025 GCTTGTATGAG         1,00         0,17 Hs.18387 transcription factor AP-2 alph enhancer           3026 CCTGTCAATGT         1,00         0,17 Hs.183745 ESTs           3028 GGGCTAGTGGG         1,00         0,17 Hs.182591 RAS guanyl releasing protein and DAG-regu           3029 GTGACGCCCCG         1,00         0,17 Hs.173993 RNA binding motif protein 6           3031 AATGTACCTGG         1,00         0,17 Hs.173422 CAMP responsive element bin protein 3 (luman)           3032 GCCCTCATTAA         1,00         0,17 Hs.169370 FYN oncogene related to SR	
3018   TAACTGTCAAA	3018 TAACTGTCAAA         1,00         0,17 Hs.238205 hypothetical protein PRO201           3019 GTGGCTGATGC         1,00         0,17 Hs.238118 EST           3020 AAATTTGTATG         1,00         0,17 Hs.235445 Homo sapiens mRNA; cDNA DKFZp761G2312 (from clone DKFZp434M0737 gene product DKFZp434M0737	jene 3
3019   GTGGCTGATGC	3019 GTGGCTGATGC	
3020   AAATTTGTATG	3020   AAATTTGTATG	
DKFZp761G2312 (from clone DK	DKFZp761G2312 (from clone	
3021   AGCCACGGC   1,00   0,17   Hs.230396   EST   3022   AGGACATTCG   1,00   0,17   Hs.214982   laminin, gamma 1 (formerly LAMB2)   3023   CATAGATTTC   1,00   0,17   Hs.194035   KIAA0737 gene product   3024   TTCTCAATACA   1,00   0,17   Hs.193164   ESTS   3025   GCTTGTATGAG   1,00   0,17   Hs.186655   Homo sapiens mRNA; cDNA   DKFZp434M0223 (from clone DK   1,00   1,0	3021 AGCCACCAGGC         1,00         0,17 Hs.230396 EST           3022 AAGGACATTCG         1,00         0,17 Hs.214982 faminin, gamma 1 (formerly L           3023 CCATAGATTTC         1,00         0,17 Hs.194035 KIAA0737 gene product           3024 TTCTCAATACA         1,00         0,17 Hs.193164 ESTs           3025 GCTTGTATGAG         1,00         0,17 Hs.186655 Homo sapiens mRNA; cDNA DKFZp434M0223 (from clone and DKFZp434M0223 (from clone contact and DKFZp434M0223 (from clone contact and DKFZp434M0223 (from clone contact and	
3022   AAGGACATTCG	3022 AAGGACATTCG         1,00         0,17 Hs.214982 Iaminin, gamma 1 (formerly L           3023 CCATAGATTTC         1,00         0,17 Hs.194035 KIAA0737 gene product           3024 TTCTCAATACA         1,00         0,17 Hs.193164 ESTs           3025 GCTTGTATGAG         1,00         0,17 Hs.186655 Homo sapiens mRNA; cDNA DKFZp434M0223 (from clone and DKFZp434M0223 (from clone contact and DKFZp434M0223 (from clone contact and DKFZp434M0223 (from clone contact and con	e DIC
3023   CCATAGATTTC   1,00	3023 CCATAGATTTC         1,00         0,17 Hs.194035 KIAA0737 gene product           3024 TTCTCAATACA         1,00         0,17 Hs.193164 ESTs           3025 GCTTGTATGAG         1,00         0,17 Hs.186655 Homo sapiens mRNA; cDNA DKFZp434M0223 (from clone and DKFZp434M0223 (from clone contact and DKFZp434M0223 (from clo	AMR2)
3024   TTCTCAATACA	3024 TTCTCAATACA         1,00         0,17 Hs.193164 ESTs           3025 GCTTGTATGAG         1,00         0,17 Hs.186655 Homo sapiens mRNA; cDNA DKFZp434M0223 (from clone DKFZp434M0223 (from cl	
3025   GCTTGTATGAG	3025 GCTTGTATGAG  1,00  0,17 Hs.186655 Homo sapiens mRNA; cDNA DKFZp434M0223 (from clone DKFZp434M0223 (from clone DKFZp434M0223 (from clone Transcription factor AP-2 alph enhancer  3027 GGAATTTGCTT  1,00  0,17 Hs.183745 ESTs  3028 GGGCTAGTGGG  1,00  0,17 Hs.182591 RAS guanyl releasing protein and DAG-regu  3029 GTGACGCCCCG  1,00  0,17 Hs.179661 tubulin, beta polypeptide  3030 CAATGAGCATC  1,00  0,17 Hs.173993 RNA binding motif protein 6  3031 AATGTACCTGG  1,00  0,17 Hs.173422 cAMP responsive element bin protein 3 (luman)  3032 GCCCTCATTAA  1,00  0,17 Hs.169370 FYN oncogene related to SR	
DKFZp434M0223 (from clone DK	DKFZp434M0223 (from clone 3026 CCTGTCAATGT 1,00 0,17 Hs.18387 transcription factor AP-2 alphenhancer 3027 GGAATTTGCTT 1,00 0,17 Hs.183745 ESTs 3028 GGGCTAGTGGG 1,00 0,17 Hs.182591 RAS guanyl releasing protein and DAG-regu 3029 GTGACGCCCG 1,00 0,17 Hs.179661 tubulin, beta polypeptide 3030 CAATGAGCATC 1,00 0,17 Hs.173993 RNA binding motif protein 6 3031 AATGTACCTGG 1,00 0,17 Hs.173422 CAMP responsive element bin protein 3 (luman) 3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	
3026   CCTGTCAATGT	3026 CCTGTCAATGT  1,00  0,17 Hs.18387 transcription factor AP-2 alphenhancer  3027 GGAATTTGCTT  1,00  0,17 Hs.183745 ESTs  3028 GGGCTAGTGGG  1,00  0,17 Hs.182591 RAS guanyl releasing protein and DAG-regu  3029 GTGACGCCCGG  1,00  0,17 Hs.173993 RNA binding motif protein 6  3030 CAATGAGCATC  1,00  0,17 Hs.173422 cAMP responsive element bin protein 3 (luman)  3032 GCCCTCATTAA  1,00  0,17 Hs.169370 FYN oncogene related to SR	
enhancer	enhancer  3027 GGAATTTGCTT 1,00 0,17 Hs.183745 ESTs  3028 GGGCTAGTGGG 1,00 0,17 Hs.182591 RAS guanyl releasing protein and DAG-regu  3029 GTGACGCCCG 1,00 0,17 Hs.179661 tubulin, beta polypeptide  3030 CAATGAGCATC 1,00 0,17 Hs.173993 RNA binding motif protein 6  3031 AATGTACCTGG 1,00 0,17 Hs.173422 CAMP responsive element bin protein 3 (luman)  3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	
3028   GGGCTAGTGGG	3028 GGGCTAGTGGG 1,00 0,17 Hs.182591 RAS guanyl releasing protein and DAG-regu 3029 GTGACGCCCG 1,00 0,17 Hs.179661 tubulin, beta polypeptide 3030 CAATGAGCATC 1,00 0,17 Hs.173993 RNA binding motif protein 6 3031 AATGTACCTGG 1,00 0,17 Hs.173422 CAMP responsive element bin protein 3 (luman) 3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	ia (activating
and DAG-regu	and DAG-regu  3029 GTGACGCCCG 1,00 0,17 Hs.179661 tubulin, beta polypeptide  3030 CAATGAGCATC 1,00 0,17 Hs.173993 RNA binding motif protein 6  3031 AATGTACCTGG 1,00 0,17 Hs.173422 CAMP responsive element bin protein 3 (luman)  3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	
3029   GTGACGCCCG   1,00   0,17   Hs.179661   tubulin, beta polypeptide	3029 GTGACGCCCG1,000,17 Hs.179661 tubulin, beta polypeptide3030 CAATGAGCATC1,000,17 Hs.173993 RNA binding motif protein 63031 AATGTACCTGG1,000,17 Hs.173422 CAMP responsive element bin protein 3 (luman)3032 GCCCTCATTAA1,000,17 Hs.169370 FYN oncogene related to SR	1 1 (calcium
3030   CAATGAGCATC   1,00   0,17   Hs.173993   RNA binding motif protein 6   3031   AATGTACCTGG   1,00   0,17   Hs.173422   CAMP responsive element binding protein 3 (luman)   3032   GCCCTCATTAA   1,00   0,17   Hs.169370   FYN oncogene related to SRC, FGR, YES   3033   ATACAAATATG   1,00   0,17   Hs.16533   myosin phosphatase, target subunit 1   3034   TTGCACTTAAT   1,00   0,17   Hs.16206   uncharacterized hypothalamus protein HT008   3035   AAAGCTGTGTT   1,00   0,17   Hs.155485   huntingtin interacting protein 2   3036   AGAATAAATCT   1,00   0,17   Hs.1554437   phosphodiesterase 2A, cGMP-stimulated 3037   GAAAACTACCC   1,00   0,17   Hs.150741   2',3'-cyclic nucleotide 3' phosphodiesterase   3038   TGGCCTAATAA   1,00   0,17   Hs.1501   syndecan 2 (heparan sulfate proteoglycan 1, cell sur 3039   GTATCAAGCAG   1,00   0,17   Hs.149436   kinesin family member 5B   3040   CCTGAGTGCGT   1,00   0,17   Hs.14595   hypothetical protein FLJ20615   3041   TAACACTGACT   1,00   0,17   Hs.13572   calcium modulating ligand 3042   TAATTCTTTTA   1,00   0,17   Hs.13572   calcium modulating ligand 3043   GCTCAAAACCT   1,00   0,17   Hs.13572   calcium modulating ligand 3045   CCTCTCCAACA   1,00   0,17   Hs.118397   AE-binding protein 1   major histocompatibility complex, class III, DM beta   3046   AGGGATATGGG   1,00   0,17   Hs.114434   ESTs, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J S   3047   GATCAAAATTT   1,00   0,17   Hs.109805   Homo sapiens cDNA FLJ20153 fis, clone   COL08656, high   3048   AGAGGGACAAC   1,00   0,17   Hs.10647   FLJ00005 protein	3030 CAATGAGCATC 1,00 0,17 Hs.173993 RNA binding motif protein 6 3031 AATGTACCTGG 1,00 0,17 Hs.173422 cAMP responsive element bin protein 3 (luman) 3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	
3031   AATGTACCTGG	3031 AATGTACCTGG 1,00 0,17 Hs.173422 cAMP responsive element bii protein 3 (luman) 3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	
Description of the state of t	protein 3 (luman) 3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	
3032   GCCTCATTAA   1,00   0,17   Hs.169370   FYN oncogene related to SRC, FGR, YES	3032 GCCCTCATTAA 1,00 0,17 Hs.169370 FYN oncogene related to SR	nding
YES   3033   ATACAAATATG   1,00   0,17   Hs.16533   myosin phosphatase, target subunit 1   3034   TTGCACTTAAT   1,00   0,17   Hs.16206   uncharacterized hypothalamus protein   HT008   3035   AAAGCTGTGTT   1,00   0,17   Hs.155485   huntingtin interacting protein 2   3036   AGAATAAATCT   1,00   0,17   Hs.154437   phosphodiesterase 2A, cGMP-stimulated 3037   GAAAACTACCC   1,00   0,17   Hs.150741   2',3'-cyclic nucleotide 3' phosphodiesterase   3038   TGGCCTAATAA   1,00   0,17   Hs.1501   syndecan 2 (heparan sulfate proteoglycan 1, cell sur   3039   GTATCAAGCAG   1,00   0,17   Hs.149436   kinesin family member 5B   3040   CCTGAGTGCGT   1,00   0,17   Hs.14595   hypothetical protein FLJ20615   3041   TAACACTGACT   1,00   0,17   Hs.13809   hypothetical protein FLJ10648   3042   TAATTCTTTTA   1,00   0,17   Hs.13572   calcium modulating ligand   3043   GCTCAAAACCT   1,00   0,17   Hs.13572   calcium modulating ligand   3044   GCACGGAAAAA   1,00   0,17   Hs.118397   AE-binding protein 1   3045   CCTCTCCAACA   1,00   0,17   Hs.1162   major histocompatibility complex, class   II, DM beta   3047   GATCAAAATTT   1,00   0,17   Hs.114434   ESTs, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J S   3047   GATCAAAATTT   1,00   0,17   Hs.10647   FLJ00005 protein		
3033 ATACAAATATG	I I IYES	C, FGR,
3034   TTGCACTTAAT		
HT008	<u> </u>	subunit 1
3035   AAAGCTGTGTT   1,00   0,17   Hs.155485   huntingtin interacting protein 2   3036   AGAATAAATCT   1,00   0,17   Hs.154437   phosphodiesterase 2A, cGMP-stimulated   3037   GAAAACTACCC   1,00   0,17   Hs.150741   2',3'-cyclic nucleotide 3'   phosphodiesterase   3038   TGGCCTAATAA   1,00   0,17   Hs.1501   syndecan 2 (heparan sulfate   proteoglycan 1, cell sur   3039   GTATCAAGCAG   1,00   0,17   Hs.149436   kinesin family member 5B   3040   CCTGAGTGCGT   1,00   0,17   Hs.14595   hypothetical protein FLJ20615   3041   TAACACTGACT   1,00   0,17   Hs.13809   hypothetical protein FLJ10648   3042   TAATTCTTTTA   1,00   0,17   Hs.13572   calcium modulating ligand   3043   GCTCAAAACCT   1,00   0,17   Hs.118397   AE-binding protein 1   3045   CCTCTCCAACA   1,00   0,17   Hs.1162   major histocompatibility complex, class   II, DM   beta   3046   AGGGGATATGGG   1,00   0,17   Hs.114434   ESTs, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J S   3047   GATCAAAATTT   1,00   0,17   Hs.10647   FLJ00005   protein   3048   AGAGGGACAAC   1,00   0,17   Hs.10647   FLJ00005   protein		us protein
3036 AGAATAAATCT         1,00         0,17 Hs.154437 phosphodiesterase 2A, cGMP-stimulated           3037 GAAAACTACCC         1,00         0,17 Hs.150741 2',3'-cyclic nucleotide 3' phosphodiesterase           3038 TGGCCTAATAA         1,00         0,17 Hs.1501 syndecan 2 (heparan sulfate proteoglycan 1, cell sur           3039 GTATCAAGCAG         1,00         0,17 Hs.149436 kinesin family member 5B           3040 CCTGAGTGCGT         1,00         0,17 Hs.14595 hypothetical protein FLJ20615           3041 TAACACTGACT         1,00         0,17 Hs.13809 hypothetical protein FLJ10648           3042 TAATTCTTTTA         1,00         0,17 Hs.13572 calcium modulating ligand           3043 GCTCAAAACCT         1,00         0,17 Hs.118397 AE-binding protein 1           3045 CCTCTCCAACA         1,00         0,17 Hs.1162 major histocompatibility complex, class II, DM beta           3046 AGGGATATGGG         1,00         0,17 Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047 GATCAAAATTT         1,00         0,17 Hs.10647 FLJ00005 protein		
3037   GAAAACTACCC	3036 AGAATAAATCT 1,00 0,17 Hs. 153463 huntingtin interacting protein	
phosphodiesterase   3038 TGGCCTAATAA   1,00   0,17 Hs.1501   syndecan 2 (heparan sulfate proteoglycan 1, cell sur   3039 GTATCAAGCAG   1,00   0,17 Hs.149436   kinesin family member 5B   3040 CCTGAGTGCGT   1,00   0,17 Hs.14595   hypothetical protein FLJ20615   3041 TAACACTGACT   1,00   0,17 Hs.13809   hypothetical protein FLJ10648   3042 TAATTCTTTTA   1,00   0,17 Hs.13572   calcium modulating ligand   3043 GCTCAAAACCT   1,00   0,17 Hs.123210 ESTs   3044 GCACGGAAAAA   1,00   0,17 Hs.118397   AE-binding protein 1   3045 CCTCTCCAACA   1,00   0,17 Hs.1162   major histocompatibility complex, class   II, DM   beta   3046 AGGGATATGGG   1,00   0,17 Hs.114434   ESTs, Weakly similar to ALU1_HUMAN   ALU SUBFAMILY J S   3047 GATCAAAATTT   1,00   0,17 Hs.109805   Homo sapiens cDNA FLJ20153 fis, clone   COL08656, high   3048 AGAGGGACAAC   1,00   0,17 Hs.10647   FLJ00005 protein	3037 GAAAACTACCC 1.00 0,17 Hs. 154437 phosphodiesterase ZA, cGWii	P-stimulated
3038         TGGCCTAATAA         1,00         0,17         Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           3039         GTATCAAGCAG         1,00         0,17         Hs.149436 kinesin family member 5B           3040         CCTGAGTGCGT         1,00         0,17         Hs.14595 hypothetical protein FLJ20615           3041         TAACACTGACT         1,00         0,17         Hs.13809 hypothetical protein FLJ10648           3042         TAATTCTTTA         1,00         0,17         Hs.13572 calcium modulating ligand           3043         GCTCAAAACCT         1,00         0,17         Hs.123210 ESTs           3044         GCACGGAAAAA         1,00         0,17         Hs.118397 AE-binding protein 1           3045         CCTCTCCAACA         1,00         0,17         Hs.1162 major histocompatibility complex, class II, DM beta           3046         AGGGATATGGG         1,00         0,17         Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047         GATCAAAATTT         1,00         0,17         Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048         AGAGGGACAAC         1,00         0,17         Hs.10647 FLJ00005 protein		
proteoglycan 1, cell sur		
3039 GTATCAAGCAG         1,00         0,17 Hs.149436 kinesin family member 5B           3040 CCTGAGTGCGT         1,00         0,17 Hs.14595 hypothetical protein FLJ20615           3041 TAACACTGACT         1,00         0,17 Hs.13809 hypothetical protein FLJ10648           3042 TAATTCTTTA         1,00         0,17 Hs.13572 calcium modulating ligand           3043 GCTCAAAACCT         1,00         0,17 Hs.123210 ESTs           3044 GCACGGAAAAA         1,00         0,17 Hs.118397 AE-binding protein 1           3045 CCTCTCCAACA         1,00         0,17 Hs.1162 major histocompatibility complex, class II, DM beta           3046 AGGGATATGGG         1,00         0,17 Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047 GATCAAAATTT         1,00         0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048 AGAGGGACAAC         1,00         0,17 Hs.10647 FLJ00005 protein		•
3040         CCTGAGTGCGT         1,00         0,17         Hs.14595         hypothetical protein FLJ20615           3041         TAACACTGACT         1,00         0,17         Hs.13809         hypothetical protein FLJ10648           3042         TAATTCTTTA         1,00         0,17         Hs.13572         calcium modulating ligand           3043         GCTCAAAACCT         1,00         0,17         Hs.123210         ESTs           3044         GCACGGAAAAA         1,00         0,17         Hs.118397         AE-binding protein 1           3045         CCTCTCCAACA         1,00         0,17         Hs.1162         major histocompatibility complex, class II, DM beta           3046         AGGGATATGGG         1,00         0,17         Hs.114434         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047         GATCAAAATTT         1,00         0,17         Hs.109805         Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048         AGAGGGACAAC         1,00         0,17         Hs.10647         FLJ00005 protein	3039 GTATCAAGCAG 1 00 0 17 Hs 149436 kinesin family member 5R	
3041 TAACACTGACT         1,00         0,17 Hs.13809         hypothetical protein FLJ10648           3042 TAATTCTTTTA         1,00         0,17 Hs.13572         calcium modulating ligand           3043 GCTCAAAACCT         1,00         0,17 Hs.123210 ESTs           3044 GCACGGAAAAA         1,00         0,17 Hs.118397 AE-binding protein 1           3045 CCTCTCCAACA         1,00         0,17 Hs.1162         major histocompatibility complex, class II, DM beta           3046 AGGGATATGGG         1,00         0,17 Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047 GATCAAAATTT         1,00         0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048 AGAGGGACAAC         1,00         0,17 Hs.10647 FLJ00005 protein		5
3042         TAATTCTTTA         1,00         0,17         Hs.13572         calcium modulating ligand           3043         GCTCAAAACCT         1,00         0,17         Hs.123210         ESTs           3044         GCACGGAAAAA         1,00         0,17         Hs.118397         AE-binding protein 1           3045         CCTCTCCAACA         1,00         0,17         Hs.1162         major histocompatibility complex, class II, DM beta           3046         AGGGATATGGG         1,00         0,17         Hs.114434         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047         GATCAAAATTT         1,00         0,17         Hs.109805         Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048         AGAGGGACAAC         1,00         0,17         Hs.10647         FLJ00005 protein		9
3043 GCTCAAAACCT         1,00         0,17 Hs.123210 ESTs           3044 GCACGGAAAAA         1,00         0,17 Hs.118397 AE-binding protein 1           3045 CCTCTCCAACA         1,00         0,17 Hs.1162 major histocompatibility complex, class II, DM beta           3046 AGGGATATGGG         1,00         0,17 Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047 GATCAAAATTT         1,00         0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048 AGAGGGACAAC         1,00         0,17 Hs.10647 FLJ00005 protein		<u></u>
3044 GCACGGAAAAA 1,00 0,17 Hs.118397 AE-binding protein 1 3045 CCTCTCCAACA 1,00 0,17 Hs.1162 major histocompatibility complex, class II, DM beta 3046 AGGGATATGGG 1,00 0,17 Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S 3047 GATCAAAATTT 1,00 0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high 3048 AGAGGGACAAC 1,00 0,17 Hs.10647 FLJ00005 protein		
3045 CCTCTCCAACA 1,00 0,17 Hs.1162 major histocompatibility complex, class II, DM beta 3046 AGGGATATGGG 1,00 0,17 Hs.114434 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S 3047 GATCAAAATTT 1,00 0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high 3048 AGAGGGACAAC 1,00 0,17 Hs.10647 FLJ00005 protein		
II, DM beta   3046 AGGGATATGGG		pley class
3046         AGGGATATGGG         1,00         0,17         Hs.114434         ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S           3047         GATCAAAATTT         1,00         0,17         Hs.109805         Homo sapiens cDNA FLJ20153 fis, clone COL08656, high           3048         AGAGGGACAAC         1,00         0,17         Hs.10647         FLJ00005 protein	1 ,	piex, Giass
ALU SUBFAMILY J S   3047 GATCAAAATTT   1,00   0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high   3048 AGAGGGACAAC   1,00   0,17 Hs.10647   FLJ00005 protein		1 HUMAN
3047 GATCAAAATTT 1,00 0,17 Hs.109805 Homo sapiens cDNA FLJ20153 fis, clone COL08656, high 3048 AGAGGGACAAC 1,00 0,17 Hs.10647 FLJ00005 protein		
COL08656, high   3048 AGAGGGACAAC   1,00   0,17 Hs.10647   FLJ00005 protein		53 fis. clone
3048 AGAGGGACAAC 1,00 0,17 Hs.10647 FLJ00005 protein		,
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to to the second	3049 CCATCCAGGCA 1,00 0,17 Hs.106260 sorting nexin 10	
	3050 TGTAACGTGGG 3,00 0,44 Hs.66762 Homo sapiens mRNA; cDNA	

					DKFZp564A026 (from clone DKF
3051	CACCTTCTGCC	3,00	0,44	Hs.25511	transforming growth factor beta 1
					induced transcript
	CTGTACAGACA	35,00			tubulin, beta, 2
3053	AAAGTTTGAGA	4,00		Hs.23581	leptin receptor gene-related protein
3054	GGCTTCCTAAT	2,00	0,33	Hs.284146	hypothetical protein DKFZp762N0610
3055	CGAGGGGGGCG	2,00	0,33	Hs.169875	thrombospondin 3
3056	CCCAAGCTAGC	67,00	2,89	Hs.76067	heat shock 27kD protein 1
3057	CATATCATTAA	28,00	1,60	Hs.119206	insulin-like growth factor binding protein
3058	CTCGGAGGCCT	4,00	0,52	Hs.279623	selenoprotein X
3059	GTAGACTTGTC	4,00	0,52	Hs.25213	hypothetical protein
3060	TCTGCAAGCAG	3,00			ESTs, Weakly similar to neural variant mena++ protei
3061	ACCCCCCGCC	22,00	1.36	Hs.277167	EST
	TGCAGTGACTG	4,00			LIM domain protein
	TGGCTCCTCCC	4,00			lymphocyte cytosolic protein 1 (L-plastin)
	CACTGTGTTGA	4,00			hypothetical protein FLJ10330
	TATTTCACCGT	6,00			Rho GTPase activating protein 1
	TCCAGGAAACT	6,00			cathepsin F
	CTGGGGGGAAG	2,00			KIAA0638 protein
	GCTTTTGGAGG	2,00			KIAA1538 protein
	TGCCTTAAACA	2,00			TAR (HIV) RNA-binding protein 1
	AACATAGGAAA	2,00	0.32	Hs 119663	CD59 antigen p18-20 (antigen identified
		_,,,,	0,02	. 10. 1 10000	by monoclona
3071	TAAGAGTGCTG	2,00	0.32	Hs.117582	CGI-43 protein
_	GCTGTAGACAA	2,00		Hs.10362	
	GCGAAACCCTA	3,00		Hs.270249	
	GCTCAGATCGG	3,00			KIAA0446 gene product
	TGGCCTCTCTG	4,00			peroxisomal long-chain acyl-coA
					thioesterase; putat
	TTCTTGAACAA	11,00			amplified in osteosarcoma
3077	GTGGCACACGC	33,00	1,67		eukaryotic translation initiation factor 3, subunit
	TGTGGCCTCCT	10,00		Hs.5258	chromosome 11 open reading frame2
3079	GTTGTGATGTT	5,00	0,58	Hs.129953	Ewing sarcoma breakpoint region 1
	GTAGCAGGTGC	6,00	0,64	Hs.8728	hypothetical protein DKFZp434G171
	ACATCCCAGAA	6,00	0,64	Hs.17377	coronin, actin-binding protein, 1C
-	TAATTTTGGAA	6,00	0,64	Hs.117582	CGI-43 protein
	GAAATTTTTGA	1,00		Hs.98069	Sec23-interacting protein p125
3084	TTCTTTGAATA	1,00	0,16	Hs.979	pyruvate dehydrogenase (lipoamide) beta
3085	ACACTAAAATG	1,00	0,16	Hs.91640	nuclear factor of kappa light polypeptide gene enhan
3086	CAACCCAGATT	1,00	0.16	Hs.9096	hypothetical protein FLJ20473
	TCTTTCGTCTG	1,00		Hs.87138	ESTs .
	ACTATCTCTAG	1,00		Hs.86347	hypothetical protein
	TTTGCTTGCAA	1,00		Hs.83938	ESTs, Moderately similar to MASP-2
		.,,,,,			[H.sapiens]

3090	TTTTGTACGCA	1,00	0,16	Hs.82116	myeloid differentiation primary response
					gene (88)
3091	AAGGAGATTAT	1,00	0,16	Hs.79402	polymerase (RNA) II (DNA directed) polypeptide C (33
3092	AGGCTATTGGA	1,00			LPS-induced TNF-alpha factor
3093	CATAAATATGC	1,00	0,16	Hs.75598	heterogeneous nuclear ribonucleoprotein A2/B1
3094	TTTATAACTAT	1,00			translin
3095	GAAAATAAACT	1,00	0,16	Hs.71252	Homo sapiens mRNA; cDNA DKFZp761C169 (from clone DKF
	TAAGGCTTAAC	1,00		Hs.69423	kallikrein 10
3097	TGGTTAGATAA	1,00		Hs.6179	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k
	CAGGCTGCTGG	1,00	0,16	Hs.60440	ESTs, Weakly similar to serin protease with IGF-bind
	GCCTGGAGTTG	1,00		Hs.57877	ESTs
	ACCCACTTTCT	1,00	0,16	Hs.5716	KIAA0310 gene product
	ACGATGGCCGA	1,00	0,16	Hs.5210	glia maturation factor, gamma
3102	TAAATTGGATA	1,00	0,16	Hs.5151	RAN binding protein 7
3103	AACTGTATACA	1,00		Hs.502	ATP-binding cassette, sub-family B (MDR/TAP), member
3104	GATTCTGAGTT	1,00	0,16	Hs.47679	Homo sapiens mRNA; cDNA DKFZp564I112 (from clone DKF
3105	TTGAGTGCAGT	1,00	0,16	Hs.44754	ESTs
3106	AGTTGCAGATA	1,00	0,16	Hs.30258	ESTs
3107	GCGCACCGCTG	1,00	0,16	Hs.29385	AFG3 (ATPase family gene 3, yeast)-like 2
3108	GCTTCACACCA	1,00	0,16	Hs.284932	Homo sapiens clone 24650 ubiquitin hydrolase mRNA, p
3109	GTGGGCCAGGA	1,00	0,16	Hs.284380	gamma-glutamyltransferase 1
3110	CCAGAAAAGAG	1,00			zinc finger protein 232
3111	CCCAACTAATT	1,00	0,16	Hs.279893	hypothetical protein FLJ20342
3112	AATCCCCATCC	1,00	0,16	Hs.262292	ESTs
3113	GTGCTCATTCT	1,00	0,16	Hs.25945	ESTs
	CTCTGCTGTCT	1,00	0,16	Hs.257135	EST
3115	TCTAAATAAAA	1,00	0,16	Hs.238814	EST
3116	ACCAACTAGAA	1,00			ESTs
	GGAGGTTGAGG	1,00		Hs.228827	
	GAGAAGAAATG	1,00			checkpoint suppressor 1
3119	CCTACTAAATG	1,00			coagulation factor C (Limulus polyphemus) homology (
	AGCTACCACGC	1,00	0,16	Hs.193965	ESTs_
3121	CTAAATCACTG	1,00	0,16	Hs.19180	Homo sapiens mRNA; cDNA DKFZp564E122 (from clone DKF
3122	TGGAAGGTAAT	1,00	0,16	Hs.18724	Homo sapiens mRNA; cDNA DKFZp564F093 (from clone DKF
3123	ATTGCTAAAAA	1,00	0.16	Hs.182470	PTD010 protein
	GCCAAGGGGTT	1,00			arginyl-tRNA synthetase
	TTTTTAAAAA	1,00			interferon gamma receptor 2 (interferon

	T	г			Igamma transd
2126	CACACATCACC	1.00	0.16	Un 17040	gamma transd
	GAGAGATGACG	1,00			ESTs
	AATAAGCCAAT	1,00			butyrophilin, subfamily 2, member A1
	CTGTGCTCTAC	1,00		Hs.169501	
3129	GCTGCCAGCAC	1,00	0,16	Hs.1578	baculoviral IAP repeat-containing 5
0400	00440407077	4.00	0.40	11 454400	(survivin)
	CCAACAGTCTT	1,00			KIAA0693 protein
	CCTCCTCTGCT	1,00		Hs.153503	
	TGAAACCCTGT	1,00		Hs.142067	
	AATTGAATAAT	1,00			prefoldin 2
-	AGAAAATGTGA	1,00		Hs.127842	
$\overline{}$	CCCTCTCTGTA	1,00		Hs.126783	
3136	GGATTCTGACT	1,00	0,16	Hs.12150	retinal short-chain
					dehydrogenase/reductase retSDR2
	TTCTAAGTGTG	1,00	0,16	<u>Hs.119488</u>	cystein-rich hydrophobic domain 2
3138	GCATTGTTTTA	1,00	0,16		fucosyltransferase 8 (alpha (1,6)
					fucosyltransferase
	GAGAAGACTGT	1,00			zinc finger protein 266
	TTACAACAGCA	1,00			tetratricopeptide repeat domain 3
	GTAGAGTAGGA	1,00			hypothetical protein
	CTAATTTTACA	1,00		Hs.10283	
3143	CTGAGGGTGGT	4,00	0,50	Hs.279761	HSPC134 protein
3144	CCACTCTGGCT	3,00	0,42	Hs.83919	glucosidase I
3145	GTGAGCAAGAC	3,00	0,42	Hs.78040	KDEL (Lys-Asp-Glu-Leu) endoplasmic
					reticulum protein
	CACTGTGTGTA	3,00	0,42	Hs.164207	ESTs
	TAGCAATCAGA	2,00	0,32	Hs.83795	interferon regulatory factor 2
3148	GATGGAATGTT	2,00	0,32	Hs.76550	Homo sapiens mRNA; cDNA
					DKFZp564B1264 (from clone DK
3149	ACTGGGTGGAA	2,00	0,32	Hs.59622	ESTs, Weakly similar to unknown
					[H.sapiens]
	CACCTAATTGA	2,00			baculoviral IAP repeat-containing 2
3151	TTCACAGTGCA	2,00			hypothetical protein FLJ10420
	GTGGACTTTTG	2,00			DKFZP434J154 protein
3153	CCTTACCTACA	2,00	0,32	Hs.184542	CGI-127 protein
3154	CATAACCTTCC	2,00	0,32	Hs.167460	splicing factor, arginine/serine-rich 3
	GCTAGGTATTT	2,00		Hs.165986	
	TTGTAAAGTAA	2,00	0,32	Hs.158688	translation initiation factor IF2
3157	CCCTTGACCCT	2,00	0,32	Hs.124009	Human DNA sequence from clone RP5-
					860F19 on chromoso
3158	CCACTAATGGA	4,00	0,50		SMT3 (suppressor of mif two 3, yeast)
					homolog 2
3159	AGCTCTGCTGC	4,00	0,50	Hs.102402	Mad4 homolog
3160	TTTGCACCTTT	21,00			connective tissue growth factor
3161	GCTTTGATGAT	9,00			epoxide hydrolase 1, microsomal
					(xenobiotic)
3162	TGTGGCCTGCA	3,00	0,42	Hs.76480	ubiquitin-like 4
3163	CGGAACACCGT	3,00			villin 2 (ezrin)
	CTAACCAGACA	10,00			capping protein (actin filament) muscle

		7			I7 line hate
2165	CTTAATCCTCA	22.00	1 22	Un 224422	Z-line, beta
	CTTAATCCTGA	22,00			amino acid transporter 2
3 100	GTCACTGCCTC	5,00	0,56	ns. 105584	ribosomal protein S6 kinase, 90kD, polypeptide 4
3167	AGAGGGTGGGA	2,00	0,31	Hs.82646	heat shock 40kD protein 1
3168	TTGTTTAATTT	2,00		Hs.75546	capping protein (actin filament) muscle Z-line, alph
3169	CTTGTAAACTG	2,00	0,31	Hs.57209	ESTs
3170	GTCAAAAAAAA	2,00	0,31	Hs.271699	polymerase (DNA directed) iota
3171	GGGGTACCCCT	2,00	0,31	Hs.187520	ESTs, Weakly similar to dJ353E16.2 [H.sapiens]
3172	GTGGCATTTGC	2,00	0,31	Hs.1244	CD9 antigen (p24)
3173	ATTGATCAATA	2,00	0,31	Hs.122489	ESTs, Weakly similar to ALUC_HUMAN IIII ALU CLASS C
3174	TGGCTAAAAAA	11,00	0,85	Hs.110048	ESTs
3175	TGCCAGAAATG	3,00			KIAA0141 gene product
3176	GCTCTGGTTCT	3,00		Hs.6127	ESTs, Weakly similar to A46010 X-linked retinopathy
3177	GCTCCTTTCAC	3,00	0,41	Hs.13233	
3178	CTCTGGGATAG	4,00	0,49	Hs.285254	ESTs
	AAGAAAGGAGT	5,00			procollagen C-endopeptidase enhancer
	ATGAACCGCAG	7,00	0.66	Hs.252259	ribosomal protein S3
	GTTCACATTAG	62,00			CD74 antigen (invariant polypeptide of major histoco
3182	GATGCTTTCTC	2,00	0,31	Hs.75852	casein kinase 1, delta
	ATGACCTGAAG	2,00			follicular lymphoma variant translocation
3184	GGAACCAGGTC	2,00	0.31	Hs.7404	ESTs
	TCCTCTACCTG	2,00			SNARE associated protein snapin
	TGAAGAGACTT	2,00	0,31	Hs.240767	Human DNA sequence from clone RP1- 12G14 on chromosom
3187	GCACCAAAAAA	2,00	0.31	Hs.23585	KIAA1078 protein
	TAACCAAATAC	2,00	0.31	Hs.201623	ESTs
	TATTCCCCACC	2,00	0.31	Hs.199316	ESTs
3190	TTCCAGCTGCT	2,00	0,31	Hs.19121	adaptor-related protein complex 2, alpha 2 subunit
	GTAGACTCTTT	2,00	0,31	Hs.181300	sel-1 (suppressor of lin-12, C.elegans)- like
	CTTCAATTCTG	2,00			Homo sapiens cDNA FLJ10174 fis, clone HEMBA1003959
	ACAGCTAACAG	2,00			KIAA0935 protein
	CTCGCTTCTCC	2,00	0,31	Hs.110953	Homo sapiens mRNA; cDNA DKFZp434A139 (from clone DKF
<u> </u>	TCATTCCACTG	1,00		Hs.99804	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
3196	TACTGATTACT	1,00	0,16	Hs.9599	solute carrier family 25, member 13 (citrin)
3197	TCAGGTACTGA	1,00	0,16	Hs.83951	Hermansky-Pudlak syndrome
3198	AAACTGTGAAT	1,00		Hs.83623	nuclear receptor subfamily 1, group I,

	<u> </u>	- T		<del></del>	member 3
3199	ACTGACTGACT	1,00	0.16	Hs.83050	phosphoinositide-3-kinase, regulatory
					subunit 4, p15
3200	TACTTGAAGGA	1,00	0,16	Hs.79067	MAD (mothers against decapentaplegic, Drosophila) ho
3201	TTTAACAGAAA	1,00	0.16	Hs.77617	nuclear antigen Sp100
	TATCAAAACAC	1,00		Hs.74420	origin recognition complex, subunit 3 (yeast homolog
2202	CTTATATCCAA	1,00	0.16	Hs.71475	hypothetical protein
	GTTATATCCAA ATGAACTGATT			Hs.6630	ESTs
	CTGACGGGTAT	1,00		Hs.63795	
		1,00			Homo sapiens mRNA; cDNA DKFZp434P232 (from clone DKF
3206	GAATATGGCTA	1,00	0,16	Hs.59594	ESTs, Weakly similar to F33G12.3 gene product [C.ele
3207	CTTGGGTCCTA	1,00	0,16	Hs.5288	Homo sapiens mRNA; cDNA
2200	CCTCCCAACTC	1.00	0.16	He 40024	DKFZp434M245 (from clone DKF
	GGTGGGAACTC GAATGGGCTGG	1,00		Hs.40834 Hs.32988	transitional epithelia response protein
3209	GAATGGGCTGG	1,00	0,16	HS.32988	ESTs, Weakly similar to KIAA0412 [H.sapiens]
3210	CAGTTTGCATC	1,00	0,16	Hs.31547	NADH dehydrogenase (ubiquinone) 1
					alpha subcomplex,
	TAGTCATTGAG	1,00		Hs.28020	KIAA0766 gene product
	ACAATAATAGA	1,00		Hs.27258	calcyclin binding protein
	CCTTTCCTACC	1,00	0,16	Hs.26731	ESTs
	TTGAGCTTATG	1,00			ESTs
3215	AGCCACTGCAT	1,00	0,16	Hs.258916	ESTs
3216	GAGCAAATGTT	1,00	0,16	Hs.24724	MFH-amplified sequences with leucine- rich tandem rep
3217	TTGGTATGAAG	1,00	0.16		KIAA0020 gene product
	TGTAAAATGGG	1,00			hypothetical protein from EUROIMAGE 2260343
3219	ATTAATAAAAA	1,00	0,16		ubiquitination factor E4B (homologous to yeast UFD2)
3220	AGAATCATTTG	1,00	0,16		EST, Moderately similar to ALU7_HUMAN_ALU SUBFAMILY
	GAATTCCAGTT	1,00	0,16	Hs.235709	ESTs
	TCTCCATCACT	1,00	0,16	Hs.226770	DKFZP566C0424 protein
	AGTATGAGGAA	1,00	0,16	Hs.211600	tumor necrosis factor, alpha-induced protein 3
3224	GTAAGTGCCCA	1,00	0.16	Hs.198491	
	TGGGAAACAAG	1,00			SBBI31 protein
	AAAACTGCCTG	1,00			tumor necrosis factor receptor
					superfamily, member 1
3227	CTAAATAAAGT	1,00	0,16	Hs.172649	
	GCCATACAGCC	1,00			Homo sapiens mRNA; cDNA
					DKFZp434C2016 (from clone DK
3229	GTCTGATATCT	1,00	0,16	Hs.14920	Homo sapiens mRNA; cDNA DKFZp564I1916 (from clone DK
3230	GGCAACAAAGC	1,00	0.16	Hs.14434	ESTs
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3231	AATGGGAGTTT	1,00	0,16		ESTs, Moderately similar to
2000		1			ALU7_HUMAN ALU SUBFAMILY
	GAAACCCAAAA	1,00		Hs.125522	
	GAAAAACCCTG	1,00		Hs.124946	
3234	AGGCCCCTTAT	1,00	0,16	Hs.124740	ESTs, Moderately similar to
					ALU4_HUMAN ALU SUBFAMILY
	AATAAATTTGC	1,00			tripeptidyl peptidase II
	TCATACCATTG	1,00			KIAA0712 gene product
3237	CCAAGGATTGG	6,00		Hs.9003	Homo sapiens cDNA FLJ20672 fis, clone KAIA4492, high
3238	AAAAGATACTA	4,00	0,49	Hs.82071	Cbp/p300-interacting transactivator, with Glu/Asp-ri
3239	GGGCCCAGGGG	4,00	0,49	Hs.3803	reticulon 2
3240	GGCAAGCCCCA	110,00	3,14	Hs.252574	ribosomal protein L10a
3241	GAGCCTGGATA	3,00	0,41	Hs.9004	chondroitin sulfate proteoglycan 4
					(melanoma-associa
3242	GAGGGTTCCAG	3,00	0,41	Hs.167835	acyl-Coenzyme A oxidase 1, palmitoyl
3243	TAGACCAGATA	3,00			KIAA0515 protein
3244	CTGCCAAGTTG	27,00	1,31	Hs.75873	zyxin
3245	CCTGCCCACCC	5,00	0,55	Hs.4909	dickkopf (Xenopus laevis) homolog 3
3246	AACACATCAGC	3,00	0,40	Hs.76253	spinocerebellar ataxia 2
					(olivopontocerebellar ataxi
3247	AGCTATTCCTC	3,00	0,40	Hs.75334	exostoses (multiple) 2
3248	ACCAGCTCCCC	3,00	0,40	Hs.22546	ESTs, Weakly similar to C561_HUMAN CYTOCHROME B561 [
3249	AAGACCGAGGG	3,00	0,40	Hs.116754	ESTs, Highly similar to MPP8_HUMAN M-PHASE PHOSPHOPR
3250	тетттететет	5,00	0,54	Hs.85015	ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN
3251	GGGGCTTCCAG	2,00	0.30	Hs.9729	KIAA0239 protein
	ATGATCTGCCT	2,00		Hs.9460	ESTs
	TTGAATTTGTT	2,00		Hs.80248	RNA-binding protein gene with multiple splicing
3254	TATTTTCTAG	2,00	0,30	Hs.7579	hypothetical protein FLJ10402
	TTATATTGCCA	2,00		Hs.74569	KIAA0842 protein
	TAGCTGTCTTT	2,00		Hs.43141	DKFZP727C091 protein
	TGGCTTCCCCA	2,00			hypothetical protein FLJ10241
	GTTCCAACAAA	2,00			neuroblastoma-amplified protein
	CGAATGTCCTT	2,00		Hs.111758	
	TGTAGCTGCAA	2,00			hypothetical protein FLJ10659
	GAGCAGGAGCC	4,00		Hs.9028	histone deacetylase 5
	AAACCAAAAA	4,00		Hs.76753	endoglin (Osler-Rendu-Weber syndrome 1)
3263	GCTTTTATTCA	4,00	0.48	Hs.31819	HT014
	AGGAACACAAA	4,00			eukaryotic translation initiation factor 2,
			•		subunit
	TCTCTTTTTCT	7,00			epididymal secretory protein (19.5kD)
	CCCCAGCCAGT	75,00			ribosomal protein S3
3267	AGTCGCCTTCA	3,00	0,40	Hs.7811 _	eukaryotic translation initiation factor 3,

	l				subunit
3269	GAGGGTCTTGT	3,00	0.40	He 256310	SH3 domain-containing protein 6511
	GCGGCTGACAG	3,00			ESTs, Moderately similar to RNA
3203	GCGGCTGACAG	3,00	0,40	13.230311	splicing-related pro
3270	ATGTGTAACGA	10,00	0.76	Hs.81256	S100 calcium-binding protein A4
3210	AIGIGIAACGA	10,00	0,70	113.01230	(calcium protein, ca
3271	GGGTTTGTTTC	5,00	0.53	Hs.75969	proline-rich protein with nuclear targeting
027	00011101110	3,00	0,00	113.7 0000	signal
3272	GCAACTTGGAG	5,00	0.53	Hs.33787	vinexin beta (SH3-containing adaptor
		0,55	0,00	1.0.007.07	molecule-1)
3273	TTGCCAACACC	2,00	0.30	Hs.80919	synaptophysin-like protein
	AATGTGAAATG	2,00		Hs.61164	ESTs, Weakly similar to unknown
		_,,,,,	0,00		[D.melanogaster]
3275	GGAATGCCTCT	2,00	0.30	Hs.43125	ESTs
	TTCTAGTCTGC	2,00			baculoviral IAP repeat-containing 2
	GTTATAATACG	2,00			serum/glucocorticoid regulated kinase
	CCAAGGGTCCA	2,00		Hs.13494	ESTs
	GGAGTCCTAGC	2,00			hypothetical protein FLJ20411
	GAACTGAAAAA	1,00		Hs.92381	nudix (nucleoside diphosphate linked
		.,	-,		moiety X)-type
3281	CTTCAAGAGTT	1,00	0.15	Hs.90093	heat shock 70kD protein 4
	TTCTGTCCCTT	1,00		Hs.89781	upstream binding transcription factor,
		.,	-,		RNA polymeras
3283	CCTACAGACAC	1,00	0.15	Hs.8904	lg superfamily protein
	CTTACATTTTA	1,00			5T4 oncofetal trophoblast glycoprotein
	TTCTGAAGACA	1,00		Hs.76901	for protein disulfide isomerase-related
	ATTCTTAAACA	1,00			tyrosine 3-monooxygenase/tryptophan 5-
	,	'	·		monooxygenase
3287	TCCTGAATGAA	1,00	0,15	Hs.72085	hypothetical protein FLJ10853
3288	TGTGTTTATTG	1,00	0,15	Hs.6809	RAP2A, member of RAS oncogene
			·		family
3289	GCAACGTAATA	1,00	0,15	Hs.65329	kraken-like
3290	CAGTAAAGGTT	1,00	0,15	Hs.6163	ESTs, Weakly similar to myotonic
					dystrophy kinase [H
3291	AAATCCTCAAA	1,00			hypothetical protein FLJ10307
	GAGCCTCACAC	1,00		Hs.51039	KIAA0076 gene product
	CCTTTCTTTAT	1,00		Hs.50535	
3294	GAAGTTCTCTG	1,00	0,15	Hs.47438	SH3 domain binding glutamic acid-rich
					protein
	AGGGCACAGGG	1,00			ESTs
	TCAATATCACT	1,00	0,15	Hs.29276	hypothetical protein FLJ20457
3297	TTTTTATTTAG	1,00	0,15	Hs.284169	Homo sapiens mRNA full length insert
					cDNA clone EURO
3298	CACAGTATTTG	1,00	0,15	Hs.279009	seven in absentia (Drosophila) homolog
					1
3299	TGAGAGTATTT	1,00	0,15	Hs.273391	Homo sapiens mRNA; cDNA
					DKFZp761N05121 (from clone D
	CAATAAAACTG	1,00			hypothetical protein FLJ10422
<u> 3301</u>	GTGGCCGGCAC	1,00	0,15	Hs.258519	ESTs, Weakly similar to ALU5_HUMAN

	<del></del>	1			ALLI OLIDEAMU V.CO
0000	1001700077	1 25	6.45	11-05555	ALU SUBFAMILY SC
	AGGATCGCTTG	1,00		Hs.255687	
	TTCCAGTAAAC	1,00			DKFZP586F1524 protein
	TTTTCCACATC	1,00		Hs.21291	
	GTGACAGAGTG	1,00			hypothetical protein PRO2738
	TCACACTGGCT	1,00			TAL1 (SCL) interrupting locus
	GGACTGGGTCG	1,00	0,15	Hs.194714	synaptosomal-associated protein, 29kD
3308	GTGAACCCCAT	1,00			membrane fatty acid (lipid) desaturase
3309	TGGCTGCATAG	1,00	0,15	Hs.164478	ESTs, Weakly similar to AF161483_1 HSPC134 [H.sapien
3310	AACTCAGCTCC	1,00	0,15	Hs.156044	ESTs
3311	GAATAAACCAT	1,00			protein tyrosine phosphatase, non- receptor type 1
	CCTGCCGCCTA	1,00			epidermal growth factor receptor substrate EPS15R
	CACACCATTGT	1,00	0,15	Hs.136644	CS box-containing WD protein
	GTTCTGTTAAA	1,00	0,15	Hs.13525	hypothetical protein
3315	TTGACAAATTG	1,00	0,15	Hs.130824	ESTs
3316	ATATACTGTAC	1,00	0,15	Hs.126857	ESTs
3317	CGCCGAGCACG	1,00			ESTs, Weakly similar to putative
			·		serine/threonine pr
3318	ACATTITAAG	1,00	0.15	Hs.117582	CGI-43 protein
	GACGGTATCAC	1,00	0.15	Hs.112110	PTD007 protein
	GAAATATTGCT	1,00	0.15	Hs.11123	ESTs, Weakly similar to B38919
		.,55	,,,,		hypothetical protein
3321	AAACTACCCTT	1,00	0.15	Hs.101282	Homo sapiens mRNA; cDNA
		,,,,,	-,		DKFZp434B102 (from clone DKF
3322	CAGCAGTAGCG	5,00	0,53	Hs.74441	chromodomain helicase DNA binding protein 4
3323	GTGGAATAAAG	3,00	0,40	Hs.83337	latent transforming growth factor beta binding prote
3324	AAGGTGGAGGA	148,00	3.27	Hs. 163593	ribosomal protein L18a
	GGTGGCACTCA	25,00			ras homolog gene family, member A
	AATCTTGCAAA	5,00	0.53	Hs.109201	CGI-86 protein
	TCAACTGAAGT	4,00			metallothionein 1E (functional)
	GTGGCGCGCAC	4,00			ESTs, Weakly similar to ALU8_HUMAN ALU SUBFAMILY SX
	CTCACCGCCCT	4,00	0,47	Hs.183650	cellular retinoic acid-binding protein 2
3330	AGATAACACAG	3,00	0,39	Hs.194369	arginine-glutamic acid dipeptide (RE) repeats
3331	CCTTGTCTTTT	3,00	0,39	Hs.183438	DKFZP566J153 protein
3332	GAGTAAAAAA	3,00		Hs.17752	phosphatidylserine-specific phospholipase A1alpha
	GCGGCTTTCCG	6,00	0,58	Hs.278431	SCO (cytochrome oxidase deficient, yeast) homolog 2
3334	GTGGTGCGCAC	6,00	0,58	Hs.11090	high affinity immunoglobulin epsilon receptor beta s
3335	AAATAAAAGCT	8,00	0,66	Hs.155191	villin 2 (ezrin)
3336	TCACCTTAGGT	11,00			integral membrane protein 2B

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3337	CTTCTAGCAAA	2,00	0,30	Hs.9042	Homo sapiens EST from clone 491476,
2220	GCAAATGCCGT	2,00	0.30	Hs.8551	full insert PRP4/STK/WD splicing factor
	TGTCCTCCCCA	2,00			Homo sapiens clone 25020 mRNA
3338	IGICCICCCA	2,00	0,30	ПS.0211 <del>3</del>	sequence
3340	CCGCTTCTGCT	2,00	0,30	Hs.5245	hypothetical protein FLJ20643
3341	TACCCTGGAAC	2,00	0,30	Hs.283048	hypothetical protein PRO0128
3342	GTCGGACACTG	2,00		Hs.278559	
3343	TCCTGTAAAGG	11,00	0,76	Hs.74034	Homo sapiens clone 24651 mRNA sequence
3344	AGGATGACCAG	4,00	0,46	Hs.69554	hypothetical protein FLJ20552
3345	TTGCAACCAAA	4,00	0,46	Hs.10101	ESTs, Weakly similar to coded for by C. elegans cDNA
3346	TGAGGGTTAGA	3,00	0,39	Hs.6603	hypothetical protein FLJ20296
	CTCCCCATCAG	3,00		Hs.238030	secretory carrier membrane protein 2
	CTGGGATCATC	5,00			ESTs
3349	CCTAAGGCTAA	5,00	0,52	Hs.108371	E2F transcription factor 4, p107/p130- binding
3350	GATTAAACCAG	6,00	0,57	Hs.90375	hypothetical protein FLJ10597
3351	GTTGTGGTTAA	115,00		Hs.75415	beta-2-microglobulin
	GCTGGCAGGCC	4,00			choline kinase-like
-	TAGGAAAGTAA	3,00		Hs.62192	coagulation factor III (thromboplastin, tissue facto
3354	AGGCCCTGCTC	3,00	0,39	Hs.6193	Homo sapiens mRNA; cDNA DKFZp434C1717 (from clone DK
3355	ACAGAGCACAG	2,00	0,29	Hs.78672	laminin, alpha 4
3356	GCACTTCAAAC	2,00	0,29	Hs.66191	Homo sapiens clone 24675 mRNA sequence
3357	CCTGCCTCGTA	2,00	0,29	Hs.61490	schwannomin interacting protein 1
	AGCCAGCCATT	2,00			nuclear factor I/B
	CTTTATGTGTA	2,00			PRO2047 protein
	CATTAAAGGGT	2,00			CTL2 gene
	GTGGCGCGTGT	2,00			KIAA1002 protein
	GAATAACAACC	1,00		Hs.99603	ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY
3363	TTTGTGAATAT	1,00	0,15	Hs.91393	Homo sapiens clone HQ0189 PRO0189 mRNA, complete cds
3364	AGAAAGGGAGA	1,00		Hs.819	homeo box B7
3365	CTTAATGGTGT	1,00		Hs.8026	ESTs, Moderately similar to p53 regulated PA26-T2 nu
3366	TTTGTCATTTT	1,00	0,15	Hs.7918	uncharacterized hypothalamus protein HSMNP1
3367	AAAGAATATGA	1,00	0,15	Hs.79162	structure specific recognition protein 1
	TAATGCTAAAA	1,00		Hs.78944	regulator of G-protein signalling 2, 24kD
3369	AATTAATTAAA	1,00	0,15	Hs.74050	follicular lymphoma variant translocation
3370	AGAAGACTGGT	1,00	0,15	Hs.71523	ESTs
	ATTCAACAATT	1,00		Hs.7120	cytokine receptor-like molecule 9
3372	CTTAAAGTCTG	1,00	0,15	Hs.7099	hypothetical protein FLJ20265

3373	GGCCCTTGCCT	1,00	0.15	Hs.6259	ESTs
	TTGGAGCAAAG	1,00		Hs.55346	ESTs, Weakly similar to Z141_HUMAN
.	1100,100,110	1,00	0, 10	110.00040	ZINC FINGER PROTE
3375	CAGAAGAGAAG	1,00	0.15	Hs.5003	KIAA0456 protein
	GTATTTATCTT	1,00		Hs.34804	ESTs
	GACTCTGGAAA	1,00	_	Hs.2953	ribosomal protein S15a
	GGGTGGGCAGT	1,00			F-box only protein 6
	AAATATAATGA	1,00			adrenal gland protein AD-004
	AAGCAAGTTCC	1,00		Hs.277076	
	CTCACAAAAAA	1,00		Hs.27356	
	TATATAGGTCC	1,00			Homo sapiens mRNA; cDNA
					DKFZp434E1723 (from clone DK
	AAATCAATAAA	1,00		Hs.270552	
3384	TTTATTATTT	1,00		Hs.23703	
3385	GGCCGTTGACC	1,00			KIAA0844 protein
	TGTTAATTTAT	1,00			KIAA0194 protein
	AAGCTGGCCCA	1,00			hypothetical protein FLJ20203
3388	ACAGCTACAGT	1,00	0,15	Hs.19631	ESTs
3389	GTGATGCACGC	1,00		Hs.193396	ESTs, Weakly similar to alternatively
					spliced produc
3390	AAGTTCCCATT	1,00	0,15	Hs.189886	ESTs
3391	ACCAGAGAGCA	1,00	0,15	Hs.181185	DKFZP564D116 protein
3392	GAAGATGAAGG	1,00			catenin (cadherin-associated protein),
					alpha 1 (102k
3393	TAAATTCAAGC	1,00	0,15	Hs.174905	KIAA0033 protein
3394	AAATGTAACAT	1,00	0,15	Hs.164649	hypothetical protein DKFZp434H247
3395	AAACAGAGCTG	1,00	0,15	Hs.161554	hypothetical protein FLJ20159
3396	GGAGCTGTCTG	1,00			Human DNA sequence from clone RP3-
					329A5 on chromosom
	ACAAAGTTGTG	1,00	0,15	Hs.154510	carbonyl reductase 3
	AGTATGCAGAG	1,00	0,15	Hs.144407	hypothetical protein FLJ10956
3399	GATGGATGGAA	1,00		Hs.137516	
3400	TGGTATTTCGG	1,00			ESTs, Weakly similar to SP49_HUMAN
					SPLICEOSOME ASSOC
	TCTAAAAGAAA	1,00		Hs.13328	
	GGAGAGGCCCC	1,00	0,15	Hs.129928	KIAA0454 protein
	AAATAAAAGAT	1,00			gastrointestinal peptide
	AACTAAAAAAC	1,00			hypothetical protein PRO1197
3405	GGTCAAGCCAG	1,00		Hs.115222	
3406	ACTCAGATGCC	1,00	0,15	Hs.11367	hypothetical protein RP1-317E23
3407	AAGTTGTGAAG	1,00			hypothetical protein LOC55565
3408	CCTGTATTTGC	1,00		Hs.103158	
	CCTTCGAGATC	52,00			ribosomal protein S5
3410	TGGTGTTGAGG	147,00			ribosomal protein S18
	AGAACCTTCCA	17,00			major histocompatibility complex, class I,
					A
3412	ACCTGTATCCC	34,00	1,21	Hs.182241	interferon induced transmembrane
					protein 3 (1-8U)
3413	GGGAGGTAGCA	3,00	0,38	Hs.171825	basic helix-loop-helix domain containing,

					class B, 2
3414	AGCAGATCAGG	67,00	1,66	Hs.119301	S100 calcium-binding protein A10 (annexin II ligand,
3415	AGTCTGATGTT	14,00	0,81	Hs.173255	small nuclear ribonucleoprotein polypeptide A
3416	GATACACTGGC	4,00	0.45	He 25797	splicing factor 3b, subunit 4, 49kD
	GGCCCCTCCCT	4,00			chromobox homolog 6
	GCCAGGGCCAC	2,00			MYC-associated zinc finger protein (purine-binding t
3419	GCTTAATGTTT	2,00	0.29	Hs.76359	catalase
	ACAGTGTGAGA	2,00			thymine-DNA glycosylase
	GGAGAGGAAGT	2,00			ESTs
	ACAACATAGAA	2,00			ESTs
	TTAAAAGTCAC	2,00	0,29	Hs.12210	Homo sapiens cDNA FLJ11289 fis, clone PLACE1009621
3424	GAGTTTGGCCC	2,00			Mad4 homolog
	CGACGAGGAGG	12,00			epithelial membrane protein 3
3426	GCTCACTGCAG	11,00		Hs.128605	
3427	ATTCCTGACCC	3,00	0,38	Hs.166204	PHD finger protein 1
3428	CCCTGATTTTA	13,00	0,77	Hs.183684	eukaryotic translation initiation factor 4 gamma, 2
3429	CCAGAGAACTT	12,00	0,74	Hs.243886	nuclear autoantigenic sperm protein (histone-binding
3430	CTATGGCTTCA	5,00	0,50	Hs.75618	RAB11A, member RAS oncogene family
3431	AGTTGTCACTT	· 3,00	0,38	Hs.74649	cytochrome c oxidase subunit VIc
3432	TGGGGGCCGAT	3,00			transducin-like enhancer of split 2, homolog of Dros
3433	TGCAAAAAAA	3,00	0,38	Hs.181624	
3434	TGGAGATGTGA	3,00	0,38	Hs.16130	Homo sapiens cDNA FLJ10064 fis, clone HEMBA1001450
3435	GCGGCAGTTAC	2,00	0,29	Hs.82201	casein kinase 2, alpha prime polypeptide
	GCCCTGTAGTT	2,00		Hs.76578	protein inhibitor of activated STAT3
_	TCACAAGCCAC	2,00		Hs.34953	ESTs
	TATGAAAACAT	2,00		Hs.3337	transmembrane 4 superfamily member 1
3439	GGAGAAACAGC	2,00		Hs.286035	
	GGGCCTGAGT	2,00	0,29	Hs.284280	hypothetical protein DKFZp547H236
	AGGCCAGGAGT	2,00			lethal (3) malignant brain tumor l(3)mbt protein (Dr
3442	AGGTCGGGAGA	2,00	0,29	Hs.193292	
	AAATGCAATAA	2,00			nuclear transcription factor Y, gamma
	CGCCTGTGGTC	2,00			Homo sapiens clone 24528 mRNA sequence
3445	ACTITIGCCCC	2,00	0.29	Hs.10117	CGI-15 protein
	ACCTAACCGTC	1,00		Hs.9965	ESTs
	TCAGTGTATTA	1,00		Hs.9853	ESTs
	CCTCTGTTTCT	1,00		Hs.91684	Homo sapiens cDNA FLJ20148 fis, clone COL08032, high
3449	TCCGTGTGTCA	1,00	0,15	Hs.83869	hypothetical protein
	TATTAAAATAG	1,00		Hs.82028	transforming growth factor, beta receptor

					II (70-80k
2/51	AGGCCGACTG	1,00	0.15	Hs.80976	antigen identified by monoclonal
					antibody Ki-67
3452	ATGCTTTCACA	1,00	0,15	Hs.80741	propionyl Coenzyme A carboxylase, alpha polypeptide
3453	TAACCTTGCTG	1,00	0.15	Hs.80562	gelsolin (amyloidosis, Finnish type)
	GGAAATAAAAA	1,00			KIAA0652 gene product
	CCTTGTTAGCA	1,00		Hs.788	A kinase (PRKA) anchor protein (gravin)
					12
	CTTTGTACACT	1,00		Hs.74649	cytochrome c oxidase subunit VIc
3457	AAATTACATAG	1,00		Hs.73291	hypothetical protein FLJ10881
3458	AAGGCCATCTT	1,00	0,15	Hs.65328	Homo sapiens cDNA FLJ10854 fis, clone NT2RP4001507
3459	AGTGCTGAGGG	1,00	0,15	Hs.6241	phosphoinositide-3-kinase, regulatory subunit, polyp
3460	GATGGGACCAT	1,00	O 15	Hs.46829	Homo sapiens mRNA full length insert
					cDNA clone EURO
3461	ATTTTGGGCAA	1,00	0,15	Hs.28827	mitogen-activated protein kinase kinase kinase
3462	TAATTTAAAAA	1,00	0.15	Hs 283682	hypothetical protein PRO0992
	GTGAGAAGAGA	1,00		Hs.274318	
	GGCAAGCCCCC	1,00			ribosomal protein L10a
	TGATGTCTGCC	1,00		Hs.252514	
	TTCTTGTTTGT	1,00		Hs.250824	
	GACCTTGATCG	1,00			integrin, alpha 6
	TTGAAATTGTA			Hs.21857	
	TATTATTTCAG	1,00		Hs.21255	
	GAGTAATGGAT				
		1,00			hypothetical protein FLJ10534
	GCAACAAATCC	1,00			cyclin-dependent kinase 2
	ACTACAGCCAT	1,00			ubiquitin specific protease 25
34/3	CCTGATCTGTA	1,00	0,15	HS.181357	laminin receptor 1 (67kD, ribosomal protein SA)
3474	CTAAAGTACTT	1,00	0,15	Hs.171553	ESTs
3475	TTCAAGTGAAA	1,00	0,15	Hs.169681	death effector domain-containing
3476	AACTGGAGTCT	1,00	0,15	Hs.169600	KIAA0826 protein
3477	ACCTGCTTAAC	1,00	0,15	Hs.169149	karyopherin alpha 1 (importin alpha 5)
3478	AAAATTGTAAC	1,00	0,15	Hs.166982	phosphatidylinositol glycan, class F
3479	GCCGGCTGTCT	1,00			HSPC141 protein
3480	CCAGCCAGGTG	1,00			KIAA0261 protein .
3481	AAACTAGAAAT	1,00	0,15	Hs.153260	Homo sapiens c-Cbl-interacting protein (CIN85) mRNA,
3482	GGGGTCGGG	1,00	0.15	Hs.147996	protein kinase, X-linked
	AATGTTTAACG	1,00		Hs.13366	
$\overline{}$	TGGTGTTTTGG	1,00		Hs.13234	
	CATCTTATGAA	1,00		Hs.12891	
	AGAGCACACCT	1,00			fibronectin 1
	AACAGGGACAG	1,00			zinc finger protein 213
	TTAAAATACAG	1,00		Hs.11356	
	AGGGCTTGGAG	1,00			arrestin, beta 1
2703	NOGGOT I GGAG	1,00	U, 10	1113.1122/0	jan coull, beta i

3490	GCCTGGACCAT	1,00	0,15	Hs.10964	Homo sapiens mRNA; cDNA DKFZp434L0816 (from clone DK
2401	TAGCAGCAACC	1,00	0.15	Ha 100070	cyclin-dependent kinase inhibitor 1C
L					(p57, Kip2)
	CCTTTCACACA	16,00			general transcription factor II, i
3493	CAGAGACGTGG	5,00	0,50	Hs.76111	dystroglycan 1 (dystrophin-associated
				L	glycoprotein 1
	TTTTCAAGAAG	4,00		Hs.75447	ralA binding protein 1
3495	TCTACTTTTGT	4,00	0,44	Hs.74598	polymerase (DNA directed), delta 2, regulatory subun
3496	TAACAGCCAGG	11,00	0,70	Hs.81328	nuclear factor of kappa light polypeptide gene enhan
3497	TCAGATCTTTG	117,00	1,95	Hs.75344	ribosomal protein S4, X-linked
	CACCAGCATTG	9,00		Hs.75847	chromosome 15 open reading frame 3
	GCATACCTGCA	3,00		Hs.8258	DKFZP434D1335 protein
	CTGCCCCCACC	3,00			chromosome 16 open reading frame 7
	GTGCAGGCTCC	3,00			ATP-binding cassette, sub-family B (MDR/TAP), member
3502	TCAGTGAACGC	5,00	0,49	Hs.78504	inner membrane protein, mitochondrial (mitofilin)
3503	GCCCCTGCGCA	4,00	0,44	Hs.267200	ESTs, Moderately similar to T20D3.3 [C.elegans]
3504	AAGAGGCTTCG	2,00	0,28	Hs.90017	ESTs, Weakly similar to Ig-like membrane protein [H.
3505	ATTTAGCAAGC	2,00	0,28	Hs.83213	fatty acid binding protein 4, adipocyte
	CACTGAGCCAA	2,00	-	Hs.7960	hypothetical protein FLJ20027
	GGATGATGTCT	2,00		Hs.74861	activated RNA polymerase II
1		,	•		transcription cofactor 4
3508	TGTTTCAGGAT	2,00	0,28	Hs.6216	tumorous imaginal discs (Drosophila) homolog
3509	TCTGTGCTGTC	2,00	0,28	Hs.231301	
	AAGAACTAAAA	2,00			hypothetical protein
3511	CCCTGCTTCCA	2,00			KIAA1306 protein
	GCTACTCTTTG	2,00			Human clone 23652 mRNA sequence
3513	CATTGAGCTCC	2,00			SnRNP assembly defective 1 homolog
	GCACTTACAAA	2,00			Homo sapiens mRNA; cDNA
1 :					DKFZp564H2416 (from clone DK
3515	TTCTGCTCTTG	5,00	0,49	Hs.110802	von Willebrand factor
3516	GCTGAACGCGT	6,00	0,53	Hs.99029	CCAAT/enhancer binding protein (C/EBP), beta
3517	GGGAAACCCCG	6,00	0.53	Hs.254283	
	CCACTTCCTCT	3,00			KIAA0242 protein
	TTCAGCGTTCT	3,00			hypothetical protein
		5,55	-,		MPMGp800B12492Q3
3520	AGCCAAAAAA	11,00	0.69	Hs.63525	poly(rC)-binding protein 2
	ATGGCCTCCTC	4,00			syntaxin 4A (placental)
	ATTTCAAGATG	4,00			carbonic anhydrase II
	GTGATGGATGG	5,00			Homo sapiens mRNA; cDNA
		'-	,		DKFZp586O1919 (from clone DK

3524	GATTTGTGTTC	6,00	0.53	Hs 173125	peptidylprolyl isomerase F (cyclophilin F)
	TTAAAAAAAAA	9,00			yeast Sec31p homolog
	AAGAAAACTGT	6,00			KIAA1522 protein
	TGATCTGCCTG	4,00		Hs.5723	ESTs, Weakly similar to CA12_HUMAN
002,	10/1/0100010	4,00	0, 10	110.0720	COLLAGEN ALPHA 1(
3528	TGCTGCATTGA	3,00	0.37	Hs.5344	adaptor-related protein complex 1,
	100100,1110,1	0,00	0,0.	0.00	gamma 1 subunit
3529	GCAGAGATGGG	3,00	0.37	Hs.39850	hypothetical protein FLJ20517
	TTTCTGGAGGT	3,00			KIAA0545 protein
	GCAGAACCATT	2,00		Hs.96264	alpha thalassemia/mental retardation
		_,	0,20		syndrome X-link
3532	CCTAGTAAAAA	2,00	0,28	Hs.96247	translin-associated factor X
	GCACAATGGGA	2,00		Hs.85838	solute carrier family 16 (monocarboxylic
					acid transp
3534	CAGTCCCCCTC	2,00	0,28	Hs.82563	KIAA0153 protein
3535	TAGACTGGCAC	2,00	0,28	Hs.7970	DKFZP434B027 protein
3536	CACTCAATAAA	2,00	0,28	Hs.79361	kallikrein 6 (neurosin, zyme)
3537	TTCAGGAGGGG	2,00	0,28	Hs.5890	ESTs, Weakly similar to A49134 lg
					kappa chain V-I re
3538	CCAGTGGCTCA	2,00	0,28	Hs.5753	inositol(myo)-1(or 4)-monophosphatase
					2
	TGTATTTTGAC	2,00			predicted osteoblast protein
	CCGATTTTTAA	2,00			hypothetical protein FLJ20422
3541	GAATTTGGGAT	2,00	0,28		ESTs, Highly similar to KIAA0780 protein
					[H.sapiens]
	AAAGTGAAAAA	2,00			glucuronidase, beta
3543	AAAGTGGGTGG	2,00	0,28		wingless-type MMTV integration site
					family, member 4
	TGACTGGCAAA	2,00	0,28	Hs.130849	ESTs
	TTACAGAGCTT	2,00		Hs.10590	zinc finger protein 313
3546	GTAACAAAATG	1,00	0,14	Hs.99437	Homo sapiens mRNA; cDNA
05.47	OTA 0000 A 0 0 0	4.00	- 0.44	11-0000	DKFZp586G1924 (from clone DK
	CTACCCAACAG	1,00		Hs.9292	ESTs
	GAAACCCCAGA	1,00			hypothetical protein
3548	AACACCCTTTC	1,00	0,14	Hs.82911	protein tyrosine phosphatase type IVA,
2550	TATAAAAGTGG	1.00	0.14	Hs.82084	member 2 integrin beta 3 binding protein (beta3-
3550	INIMMAGIGG	1,00	Ų, 14	ПS.02U04	endonexin)
3551	GCACCCACTGG	1,00	0.14	Hs.8184	ESTs
	GAAACCTGAAA	1,00		Hs.75772	nuclear receptor subfamily 3, group C,
0002	0,440010/44	1,00	U, 14	10.10112	member 1
3553	TGTGGGAGTAG	1,00	0.14	Hs.6783	ESTs
	TGCTCCCTTTA	1,00		Hs.65450	reticulon 4
	CTGGTCGTTGG	1,00		Hs.5985	Homo sapiens clone 25186 mRNA
		',""	-,		sequence
3556	AGATTGCTGTT	1,00	0.14	Hs.59838	hypothetical protein FLJ10808
	GCCTGGCACCG	1,00		Hs.58167	zinc finger protein 282
	GATCTTTTGTC	1,00		Hs.5148	FLN29 gene product
	CATTITATITC	1,00		Hs.49136	ESTs

S580   ATGTTTCTICC						
3562   ATGAGTTTCTG	3560	ATGTTTCTTCC	1,00	0,14	Hs.41143	KIAA0581 protein
3563   TTGCCCAGACT	3561	CCCTTGCACTC	1,00	0,14	Hs.284528	EST
S584   GTTTATGTTCC			1,00			
3565   CTGAACTCTTC	3563	TTGCCCAGACT	1,00	0,14	Hs.244245	EST
3566   CCATATGATCA						
3567   AGAAAAATAA	3565	CTGAACTCTTC				
3568   AGGTCAATGAA	3566	CCATATGATCA			Hs.192966	KIAA0265 protein
3569   AAACAATACAC	3567	AGAAAAAATAA	1,00	0,14		
1 mRNA, partial   3570   TACAGATCACA   1,00   0,14   Hs.173859   frizzled (Drosophila) homolog 7   3571   CTGTAAAAAA   1,00   0,14   Hs.17364   zinc finger protein 79 (pT7)   3572   GTGAAGCCCTA   1,00   0,14   Hs.171501   ubiquitin specific protease 11   3573   GTTTGGGATGA   1,00   0,14   Hs.16193   Homo sapiens mRNA; cDNA   DKFZp586B211 (from clone DKF   3574   GCCTGTGGATG   1,00   0,14   Hs.150601   chymotrypsin-like   3575   CCTCTGCCGGG   1,00   0,14   Hs.128512   ESTS   3576   TACAGTAAAAC   1,00   0,14   Hs.125819   putative dimethyladenosine transferas   3577   TCCATTGCTGG   1,00   0,14   Hs.12303   suppressor of Ty (S.cerevisiae) 6   homolog   3578   ATCGTGCCATT   1,00   0,14   Hs.110347   REV1 protein   3580   ACCTGACACCT   1,00   0,14   Hs.10881   WD repeat domain 7   3581   GAAAGTGGCTG   1,00   0,14   Hs.10881   WD repeat domain 7   3581   GAAAGTGGCTG   1,00   0,14   Hs.108660   ATP-binding cassette, sub-family C (CFTR/MRP), membe   3583   CTGCAAAGGAG   1,00   0,14   Hs.104519   phospholipase D2   3584   ATGGGCTTGAT   5,00   0,48   Hs.8185   CGI-44 protein; sulfide dehydrogenas   like (yeast)   3586   AGCTACCGGGC   4,00   0,43   Hs.9589   ubiquilin 1   3586   AGCTACCGGGC   4,00   0,43   Hs.9589   ubiquilin 1   3586   AGCTACCGGGC   4,00   0,43   Hs.9589   ubiquilin 1   3588   TGAGCCCGGCC   9,00   0,63   Hs.238839   latent transforming growth factor beta   binding prote   3589   CTCTGTAATTT   3,00   0,37   Hs.238839   latent transforming growth factor beta   binding prote   3589   CTCTGTAATTT   3,00   0,37   Hs.284181   hypothetical protein DKFZp434P0531   membrane-associated tyrosine- and threonine-specific   3591   AGGGGGGAGGG   3,00   0,37   Hs.284181   hypothetical protein DKFZp434P0531   membrane-associated tyrosine- and threonine-specific   3594   TCAGTGAACTG   2,00   0,28   Hs.6657   Novel human gene mapping to	3568	AGGTCAATGAA	1,00	0,14	Hs.18376	KIAA1319 protein
3570   TACAGATCACA   1,00   0,14   Hs.173859   frizzled (Drosophila) homolog 7   3571   CTGTAAAAAAA   1,00   0,14   Hs.17364   zinc finger protein 79 (pT7)   3572   GTGAAGCCTA   1,00   0,14   Hs.171501   ubiquitin specific protease 11   3573   GTTTGGGATGA   1,00   0,14   Hs.16193   Homo sapiens mRNA; cDNA   DKFZp586B211 (from clone DKF   3574   GCCTGTGCGGG   1,00   0,14   Hs.150601   chymotrypsin-like   3575   CCTCTGCCGGG   1,00   0,14   Hs.128512   ESTs   3576   TACAGTAAAAC   1,00   0,14   Hs.12303   suppressor of Ty (S.cerevisiae) 6   homolog   3578   ATCGTGCCATT   1,00   0,14   Hs.110630   Human BRCA2 region, mRNA sequel CG006   S579   GCGACAAAAAG   1,00   0,14   Hs.110630   Human BRCA2 region, mRNA sequel CG006   S579   GCGACAAAAAG   1,00   0,14   Hs.108378   ESTs, Weakly similar to zeste   [D.melanogaster]   S580   ACGTGACACCT   1,00   0,14   Hs.10888   WD repeat domain 7   3581   GAAAGTGGCTG   1,00   0,14   Hs.108788   ESTs, Weakly similar to zeste   [D.melanogaster]   S582   TTTGCTGAACA   1,00   0,14   Hs.10860   ATP-binding cassette, sub-family C   CFTR/MRP), membe   S583   CTGCAAAGGAG   1,00   0,14   Hs.104519   phospholipase D2   S584   ATGGGCTTGAT   5,00   0,48   Hs.8185   CGI-44 protein; sulfide dehydrogenas   like (yeast)   S587   CCAGGCACGCT   7,00   0,43   Hs.9589   ubiquilin 1   3586   AGCTACCGGCC   4,00   0,43   Hs.9589   ubiquilin 1   3586   CCAGGCACGCT   7,00   0,56   Hs.198427   hexokinase 2   3588   TGAGCCCGGCC   9,00   0,63   Hs.238839   latent transforming growth factor beta   binding prote   3590   CTCAGCCTGAG   3,00   0,37   Hs.284181   hypothetical protein DKFZp434P0531   membrane-associated tyrosine- and threonine-specific   Cytokine receptor-like molecule 9   3594   TCAGTGAACTG   2,00   0,28   Hs.77783   membrane-associated tyrosine- and threonine-specific   Cytokine receptor-like molecule 9   3594   TCAGTGAACTG   2,00   0,28   Hs.6657   Novel human gene mapping to	3569	AAACAATACAC	1,00	0,14	Hs.181466	Homo sapiens TRAF4 associated factor
3571   CTGTAAAAAAA					L	
3572   GTGAAGCCCTA						
3573   GTTTGGGATGA						
DKFZp586B211 (from clone DKF   3574   GCCTGTGGATG   1,00   0,14   Hs.150601   chymotrypsin-like   3575   CCTCTGCCGGG   1,00   0,14   Hs.128512   ESTs   3576   TACAGTAAAAC   1,00   0,14   Hs.128519   putative dimethyladenosine transferasic   3577   TCCATTGCTGG   1,00   0,14   Hs.12303   suppressor of Ty (S.cerevisiae) 6   homolog   3578   ATCGTGCCATT   1,00   0,14   Hs.110630   Human BRCA2 region, mRNA sequel   CG006   S579   GCGACAAAAAG   1,00   0,14   Hs.10831   WD repeat domain 7   3581   GAAAGTGGCTG   1,00   0,14   Hs.10881   WD repeat domain 7   3581   GAAAGTGGCTG   1,00   0,14   Hs.108788   ESTs, Weakly similar to zeste   [D.melanogaster]   3582   TTTGCTGAACA   1,00   0,14   Hs.108660   ATP-binding cassette, sub-family C   (CFTR/MRP), membe   3583   CTGCAAAGGAG   1,00   0,14   Hs.104519   phospholipase D2   3584   ATGGGCTTGAT   5,00   0,48   Hs.8185   CGI-44 protein; sulfide dehydrogenas   like (yeast)   3585   TGTCTTTGCTC   4,00   0,43   Hs.9589   ubiquilin 1   3586   AGCTACCGGGC   4,00   0,43   Hs.6059   EGF-containing fibulin-like extracellulimatrix pro   3587   CCAGGCACGCT   7,00   0,56   Hs.198427   hexokinase 2   3588   TGAGCCCGGCC   9,00   0,63   Hs.238839   latent transforming growth factor beta   binding prote   3590   CTCAGCCTGAG   3,00   0,37   Hs.75618   RAB11A, member RAS oncogene fan   3590   CTCAGCCTGAG   3,00   0,37   Hs.284181   hypothetical protein DKFZp434P0531   membrane-associated tyrosine- and threonine-specific   3593   TGCAAAAAAA   2,00   0,28   Hs.7120   cytokine receptor-like molecule 9   3594   TCAGTGAACTG   2,00   0,28   Hs.6657   Novel human gene mapping to	-					
3574   GCCTGTGGATG   1,00   0,14   Hs.150601   Chymotrypsin-like   3575   CCTCTGCCGGG   1,00   0,14   Hs.128512   ESTs   3576   TACAGTAAAAC   1,00   0,14   Hs.128519   putative dimethyladenosine transferation   3577   TCCATTGCTGG   1,00   0,14   Hs.12303   suppressor of Ty (S.cerevisiae) 6   homolog   3578   ATCGTGCCATT   1,00   0,14   Hs.110630   Human BRCA2 region, mRNA sequel   CG006   GG066   GG06	3573	GTTTGGGATGA	1,00	0,14	Hs.16193	
3575   CCTCTGCCGGG						DKFZp586B211 (from clone DKF
3576   TACAGTAAAAC				0,14	Hs.150601	chymotrypsin-like
3577   TCCATTGCTGG						
homolog				0,14	Hs.125819	putative dimethyladenosine transferase
3578 ATCGTGCCATT	3577	TCCATTGCTGG	1,00	0,14	Hs.12303	
CG006					<u> </u>	
3579   GCGACAAAAAG	3578	ATCGTGCCATT	1,00	0,14	Hs.110630	
3580   ACGTGACACCT   1,00   0,14   Hs.10881   WD repeat domain 7						
3581 GAAAGTGGCTG 1,00 0,14 Hs.108788 ESTs, Weakly similar to zeste [D.melanogaster] 3582 TTTGCTGAACA 1,00 0,14 Hs.108660 ATP-binding cassette, sub-family C (CFTR/MRP), membe 3583 CTGCAAAGGAG 1,00 0,14 Hs.104519 phospholipase D2 3584 ATGGGCTTGAT 5,00 0,48 Hs.8185 CGI-44 protein; sulfide dehydrogenas like (yeast) 3585 TGTCTTTGCTC 4,00 0,43 Hs.9589 ubiquilin 1 3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellul matrix pro 3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2 3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote 3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fan 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs 3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3594 TCAGTGAACTG 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9						
[D.melanogaster]  3582 TTTGCTGAACA 1,00 0,14 Hs.108660 ATP-binding cassette, sub-family C (CFTR/MRP), membe  3583 CTGCAAAGGAG 1,00 0,14 Hs.104519 phospholipase D2  3584 ATGGGCTTGAT 5,00 0,48 Hs.8185 CGI-44 protein; sulfide dehydrogenas like (yeast)  3585 TGTCTTTGCTC 4,00 0,43 Hs.9589 ubiquilin 1  3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellul matrix pro  3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2  3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote  3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fand 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs  3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 membrane-associated tyrosine- and threonine-specific  3593 TGTCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific						
TTTGCTGAACA 1,00 0,14 Hs.108660 ATP-binding cassette, sub-family C (CFTR/MRP), membe  3583 CTGCAAAGGAG 1,00 0,14 Hs.104519 phospholipase D2  3584 ATGGGCTTGAT 5,00 0,48 Hs.8185 CGI-44 protein; sulfide dehydrogenas like (yeast)  3585 TGTCTTTGCTC 4,00 0,43 Hs.9589 ubiquilin 1  3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellul matrix pro  3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2  3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote  3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fand 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs  3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	3581	GAAAGTGGCTG	1,00	0,14	Hs.108788	
CFTR/MRP), membe   3583 CTGCAAAGGAG   1,00   0,14   Hs.104519   phospholipase D2   3584   ATGGGCTTGAT   5,00   0,48   Hs.8185   CGI-44 protein; sulfide dehydrogenas like (yeast)   3585 TGTCTTTGCTC   4,00   0,43   Hs.9589   ubiquilin 1   3586   AGCTACCGGGC   4,00   0,43   Hs.6059   EGF-containing fibulin-like extracellulimatrix pro   3587 CCAGGCACGCT   7,00   0,56   Hs.198427   hexokinase 2   3588 TGAGCCCGGCC   9,00   0,63   Hs.238839   latent transforming growth factor beta binding prote   3589 CTCTGTAATTT   3,00   0,37   Hs.75618   RAB11A, member RAS oncogene fant   3590 CTCAGCCTGAG   3,00   0,37   Hs.3496   ESTs   3591   AGGGGGAGGG   3,00   0,37   Hs.284181   hypothetical protein DKFZp434P0531   3592 TGGCAAAAAAA   2,00   0,28   Hs.77783   membrane-associated tyrosine- and threonine-specific   3593 TGTCAAAAAAA   2,00   0,28   Hs.7120   Cytokine receptor-like molecule 9   3594 TCAGTGAACTG   2,00   0,28   Hs.6657   Novel human gene mapping to	0500	TTTOOTOAAGA	4.00	0.44	11 100000	
3583 CTGCAAAGGAG 1,00 0,14 Hs.104519 phospholipase D2 3584 ATGGGCTTGAT 5,00 0,48 Hs.8185 CGI-44 protein; sulfide dehydrogenas like (yeast) 3585 TGTCTTTGCTC 4,00 0,43 Hs.9589 ubiquilin 1 3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellulimatrix pro 3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2 3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote 3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fan 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs 3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 3592 TGGCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	3582	ITTIGCTGAACA	1,00	0,14	Hs.108660	
3584 ATGGGCTTGAT 5,00 0,48 Hs.8185 CGI-44 protein; sulfide dehydrogenas like (yeast)  3585 TGTCTTTGCTC 4,00 0,43 Hs.9589 ubiquilin 1  3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellulimatrix pro  3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2  3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote  3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene famous and series and se	2502	CTCCAAACCAC	4.00	0.44	11- 404540	
like (yeast)						
3585 TGTCTTTGCTC 4,00 0,43 Hs.9589 ubiquilin 1 3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellul matrix pro 3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2 3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote 3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fan 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs 3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	3304	AIGGGCIIGAI	5,00	0,48	MS.8185	
3586 AGCTACCGGGC 4,00 0,43 Hs.6059 EGF-containing fibulin-like extracellul matrix pro 3587 CCAGGCACGCT 7,00 0,56 Hs.198427 hexokinase 2 3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote 3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fan 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs 3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 3592 TGGCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	2505	TOTOTTTOOTO	4.00	0.42	LIS OF SO	
matrix pro   3587 CCAGGCACGCT   7,00   0,56 Hs.198427 hexokinase 2   3588 TGAGCCCGGCC   9,00   0,63 Hs.238839 latent transforming growth factor beta binding prote   3589 CTCTGTAATTT   3,00   0,37 Hs.75618   RAB11A, member RAS oncogene fan   3590 CTCAGCCTGAG   3,00   0,37 Hs.3496   ESTs   3591 AGGGGGAGGG   3,00   0,37 Hs.284181 hypothetical protein DKFZp434P0531   3592 TGGCAAAAAAA   2,00   0,28 Hs.77783   membrane-associated tyrosine- and threonine-specific   3593 TGTCAAAAAAA   2,00   0,28 Hs.7120   cytokine receptor-like molecule 9   3594 TCAGTGAACTG   2,00   0,28 Hs.6657   Novel human gene mapping to						
3587 CCAGGCACGCT7,000,56 Hs.198427 hexokinase 23588 TGAGCCCGGCC9,000,63 Hs.238839 latent transforming growth factor beta binding prote3589 CTCTGTAATTT3,000,37 Hs.75618 RAB11A, member RAS oncogene fan3590 CTCAGCCTGAG3,000,37 Hs.3496 ESTs3591 AGGGGGAGGG3,000,37 Hs.284181 hypothetical protein DKFZp434P05313592 TGGCAAAAAAA2,000,28 Hs.77783 membrane-associated tyrosine- and threonine-specific3593 TGTCAAAAAAA2,000,28 Hs.7120 cytokine receptor-like molecule 93594 TCAGTGAACTG2,000,28 Hs.6657 Novel human gene mapping to	3300	AGCIACCGGGC	4,00	0,43	ns.6059	
3588 TGAGCCCGGCC 9,00 0,63 Hs.238839 latent transforming growth factor beta binding prote 3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fan 3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs 3591 AGGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 3592 TGGCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	3597	CCAGGCACGCT	7.00	0.56	Un 109427	
binding prote  3589 CTCTGTAATTT 3,00 0,37 Hs.75618 RAB11A, member RAS oncogene fan  3590 CTCAGCCTGAG 3,00 0,37 Hs.3496 ESTs  3591 AGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531  3592 TGGCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific  3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9  3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	3588	TGAGCCCGGCC	9.00	0,50	Ha 220020	Intent transferming grouth factor beta
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3591 AGGGGGAGGG 3,00 0,37 Hs.284181 hypothetical protein DKFZp434P0531 3592 TGGCAAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	-					
3592 TGGCAAAAAA 2,00 0,28 Hs.77783 membrane-associated tyrosine- and threonine-specific 3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9 3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to						
threonine-specific  3593 TGTCAAAAAAA 2,00 0,28 Hs.7120 cytokine receptor-like molecule 9  3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to						
3593 TGTCAAAAAA   2,00   0,28 Hs.7120   cytokine receptor-like molecule 9   3594 TCAGTGAACTG   2,00   0,28 Hs.6657   Novel human gene mapping to	0002		۷,00	0,20	10.77703	
3594 TCAGTGAACTG 2,00 0,28 Hs.6657 Novel human gene mapping to	3593	TGTCAAAAAA	2.00	0.28	Hs.7120	
chomosome 22			_,00	5,20	13.330.	
3595 GCTCAAACTAC 2,00 0,28 Hs.284332 ESTs	3595	GCTCAAACTAC	2.00	0.28	Hs.284332	
						(Manual assignment) MEMOREC PSL4
presenilin-like pro	1 1		.,	.,		

0507	00007004000	0.00	0.00	11- 400000	1
3597	GGGCTGGACGG	2,00	0,28	HS.180338	tumor necrosis factor receptor
2500	CACCCCCACAT	2.00	0.20	Un 454690	superfamily, member 1
	CAGGGCGAGAT	2,00			DKFZP434M154 protein
	GCTATCTCAGC	2,00			hypothetical SBBI03 protein
	GGAGACAGAGT	2,00	·		calmodulin 3 (phosphorylase kinase, delta)
	CAGTTGGTTGT	7,00			E1B-55kDa-associated protein 5
3602	TCAGACAAAAG	5,00	0,48	Hs.66881	Homo sapiens mRNA; cDNA DKFZp434A1518 (from clone DK
3603	CTCATTCAGCT	6,00	0,52	Hs.180139	SMT3 (suppressor of mif two 3, yeast) homolog 2
3604	GTGAAACCCTT	3,00	0,37	Hs.206955	ESTs
3605	TTCTGGACCCA	3,00	0,37	Hs.155543	proteasome (prosome, macropain) 26S subunit, non-ATP
3606	GCTTGGATCTC	12,00	0,69		FK506 binding protein 12-rapamycin associated protei
3607	CTCAACAGCAA	10,00	0,64	Hs.7811	eukaryotic translation initiation factor 3, subunit
3608	GACTCTGAAAA	6,00	0,52	Hs.2953	ribosomal protein S15a
3609	TTGCCGGTTAA	5,00	0,48	Hs.75925	proteasome (prosome, macropain) inhibitor subunit 1
3610	ACCCTGCCAAA	5,00	0,48	Hs.284546	EST
3611	TACAGAGGGAA	12,00			zinc finger protein 216
	GGGTGGGGTTG	7,00		Hs.75216	protein tyrosine phosphatase, receptor type, F
3613	GAAATACAGTT	43,00	1,07		cathepsin D (lysosomal aspartyl protease)
3614	CAACATTCCTG	11,00	0,67	Hs.180015	D-dopachrome tautomerase
3615	TCCTTGCTTCT	6,00			hypothetical protein FLJ20297
3616	CGAGGGGCCAG	33,00			actinin, alpha 4
3617	AGAGGTGGTGT	4,00	0,43	Hs.6968	KIAA1460 protein
3618	CCACGTCCATC	2,00	0,28	Hs.9018	exostoses (multiple)-like 3
3619	GCCTTGGGGGC	2,00	0,28	Hs.75658	phosphorylase, glycogen; brain
3620	AGGAGCCGGGG	2,00	0,28	Hs.25700	Homo sapiens mRNA; cDNA DKFZp434M0435 (from clone DK
3621	CGGCAAAAAAA	2,00	0,28	Hs.179747	ecotropic viral integration site 5
3622	CTGGACTCCGC	2,00	0,28	Hs.173159	transforming, acidic coiled-coil containing protein
3623	CCCGCTCTTGA	2,00	0,28	Hs.118282	
3624	GTGATATCCAA	2,00			Homo sapiens clone 25007 mRNA sequence
3625	TCAGTAAAAAT	1,00	0,14	Hs.9805	KIAA1291 protein
3626	GCAGCTATGAG	1,00		Hs.90960	ESTs
3627	CTCTCTGTGGA	1,00		Hs.8858	bromodomain adjacent to zinc finger domain, 1A
3628	TCTCAGATGAG	1,00	0,14	Hs.82568	cytochrome P450, subfamily XXVIIA (steroid 27-hydrox
3629	CTGTAGTTGCC	1,00	0,14	Hs.75798	hypothetical protein
	ATTTCACATTT	1,00		Hs.7378	Homo sapiens mRNA; cDNA

					DKFZp434G227 (from clone DKF
3631	ACTATTTCACA	1,00	0 14	Hs.72134	KIAA1064 protein
	TTGTTGAAAGG	1,00		Hs.70337	immunoglobulin superfamily, member 4
	TGCCCTGTTCC	1,00		Hs.6651	Homo sapiens clone 23645 mRNA
3033		1,00	0, 14	1 15.0051	sequence
3634	ATGATTCCCTG	1,00	0.14	Hs.64595	DKFZP566E2346 protein
	ATAATTTTTG	1,00		Hs.539	ribosomal protein S29
	GCCCACTTCCT	1,00			ESTs, Weakly similar to unknown
3030	GCCCACTICCT	1,00	0, 14	IDS.32117	
2627	CCACTCATCCC	4.00	0.14	Un 200402	[D.melanogaster]
	GGACTCATCCC CTGCCTTATTT	1,00	0,14	Hs.286102 Hs.28501	EO15
		1,00			
3039	GACAGTGGAGA	1,00	0,14	HS.284284	ESTs, Highly similar to beta-1,3-N-
2040	CACCAATAAAA	4.00	0.44	11- 004054	acetylglucosaminy
	CAGCAATAAAA	1,00	0,14	HS.284251	KIAA0544 protein
	GTGGCTCAAGC	1,00			hypothetical protein PRO0758
	ACCTTGGATTT	1,00			PRO1510 protein
	GGGCTGCTGCC	1,00		Hs.277630	
	ATGTGGGCTCA	1,00		Hs.27018	
	GGCACCGTGGC	1,00		Hs.256144	
	GAGAAACCCAG	1,00		Hs.252895	
	ATTTTGGCCAC	1,00		Hs.24831	
	TGAAGATAGAC	1,00			hypothetical protein FLJ10826
	AGCCCTTCCTC	1,00			peptide transporter 3
	CTTCTCTGTTT	1,00			insulin receptor tyrosine kinase substrate
	TTTATCCCAAA	1,00			DKFZP547E1010 protein
	CTTGTGGTCCC	1,00			interleukin 10
3653	TGCAATTCCTT	1,00			ribosomal protein L39
3654	GATATTTTTC	1,00			SH3-domain binding protein 4
3655	CTTTGTGAACA	1,00	0,14	Hs.169946	GATA-binding protein 3
3656	TAGACCCCTTG	1,00	0,14	Hs.169476	glyceraldehyde-3-phosphate
	_				dehydrogenase
3657	AGCGTGGCTCT	1,00	0,14	Hs.15589	PPAR binding protein
3658	TTTGACAATAC	1,00			BCL2/adenovirus E1B 19kD-interacting
			·		protein 2
3659	GTCTAGTCAAT	1,00	0,14	Hs.152629	KIAA0179 protein
3660	CTGGACTTTAT	1,00	0,14	Hs.146861	hypothetical protein FLJ20580
3661	TGTACTGGCAC	1,00		Hs.144090	
3662	TGAAAACCTGA	1,00			protein kinase C, nu
	GTTATATGCCC	1,00			Homo sapiens mRNA; cDNA
			•		DKFZp586D0918 (from clone DK
3664	TTAATAAATGT	1,00	0.14	Hs.13313	cAMP responsive element binding
		,		•	protein-like 2
3665	GACCAGCGGCT	1,00	0.14	Hs.127401	DKFZP434A163 protein; selective LIM
		,,,,,			binding factor,
3666	AGCCAAATAAA	1,00	0.14	Hs.123652	
	CCAACCCATTT	1,00			ESTs, Weakly similar to S64705
		.,,,,,	,		cyclophilin-like prot
3668	TTGGGGACGGG	1,00	0.14	Hs.118400	singed (Drosophila)-like (sea urchin
		.,55	-,		fascin homolog
<u> </u>	<u> </u>			<u> </u>	1

3669	TATTGACAACA	3,00	0,36	Hs.75608	tight junction protein 2 (zona occludens
3670	CAGATTAGTTA	3,00	0.36	Us 286105	Homo sapiens clone 25244 DEAD-box
3070	CAGATTAGTTA	3,00			protein p72 mRNA s
3671	AGGAGTCGACA	3,00	0,36	Hs.181369	ubiquitin fusion degradation 1-like
3672	CAAAATCAGGA	14,00	0,72	Hs.79933	cyclin I
3673	CTCTAAGAAGC	8,00	0,58	Hs.9641	complement component 1, q
					subcomponent, alpha polype
	AAACCAGGGCC	4,00	0,43	Hs.279836	HSPC166 protein
3675	ATGGCGCAGTC	4,00		Hs.239509	
3676	GGATTTGGCCT	168,00	1,70	Hs.251247	(Manual assignment) Acidic ribosomal phosphoprotein
3677	CCCCCAGATGA	6,00	0,52	Hs.25817	hypothetical protein FLJ20386
3678	TGAAAAAAAA	15,00	0,74	Hs.57904	mago-nashi (Drosophila) homolog,
			·		proliferation-assoc
3679	AAGGTGGAGTG	3,00	0,36	Hs.9573	ATP-binding cassette, sub-family F
					(GCN20), member 1
3680	TTGTTGGATAT	3,00	0,36	Hs.4099	nardilysin (N-arginine dibasic
					convertase)
	AAGTTTGCCTG	3,00		Hs.28988	glutaredoxin (thioltransferase)
	CCATTGCATTC	3,00		Hs.185156	
	TGCTGCTGCTT	5,00			hypothetical protein FLJ20396
3684	GACCCCAAGGC	14,00	0,72	Hs.82932	cyclin D1 (PRAD1: parathyroid
	=				adenomatosis 1)
	TGGTTTTGGCA	4,00			profilin 1
	GTGTCGGCTGT	4,00			eukaryotic translation elongation factor 1 beta 2
	GGCTGTACCCA	21,00	0,82	Hs.108080	cysteine and glycine-rich protein 1
3688	GTTTTTCATTG	15,00	0,73	Hs.119502	ubiquitin A-52 residue ribosomal protein fusion prod
	CGACCCTCTCC	2,00	0,28		cytochrome c oxidase subunit VIc
_	GCCACGTTGTC	2,00		Hs.32352	hypothetical protein DKFZp434K1210
	ATGTATGGGGA	2,00	0,28	Hs.283429	SMC (mouse) homolog, X chromosome
-	TGCCACCACGC	2,00		Hs.233480	EST
	CGAAAAAAAA	2,00	0,28	Hs.20815	erythroblast macrophage protein
	GTCTTCAAAGA	2,00	0,28	Hs.184227	F-box only protein 21
3695	AAGTAGAGCAG	2,00	0,28	Hs.144645	POM (POM121 rat homolog) and ZP3 fusion protein
3696	TAGAAAGGCAG	9,00	0,61		butyrate response factor 2 (EGF- response factor 2)
3697	CAGGCCCCACC	16,00	0,75	Hs.256290	S100 calcium-binding protein A11
					(calgizzarin)
3698	ATTCTGTTGTA	5,00	0,47	Hs.150580	putative translation initiation factor
3699	CGCTGGTTCCA	83,00	1,25	Hs.179943	ribosomal protein L11
	TACCCCACCTT	4,00			H2A histone family, member Y
	TGATGTTCCAC	4,00			bromodomain adjacent to zinc finger domain, 2A
3702	GAGTCTGTTCG	3,00	0,36	Hs.283636	Homo sapiens HSPC253 mRNA, partial cds
-					

3702	TTCCGGAGCCC	3 00	0.36	He 100605	hypothetical protein
	TTGCGGAGCCC	3,00	0,30	ПS. 199095	hypothetical protein
3/04	AGTCTCCCCTA	3,00	0,36	ITS. 12303	suppressor of Ty (S.cerevisiae) 6 homolog
3705	GTTGGGAGTCC	3,00	0.36	He 100504	hypothetical protein FLJ20113
	CCAGAACAGAC	119,00			ribosomal protein L30
	TAGGGCAATCT	17,00			SMT3 (suppressor of mif two 3, yeast)
ا′′۰′ا	1710000000101	17,00	0,70		homolog 2
3708	GACTTCACTTT	5,00	0 47		transferrin receptor (p90, CD71)
	GTGGACCCTGT	5,00			CGI-25 protein
	CACGCAATGCT	22,00			amino-terminal enhancer of split
	GCCTATGGTCC	8,00			HSPC141 protein
	TGGAACTGTAA	5,00			vacuolar sorting protein 4
	AACCAAAAAA	5,00			isocitrate dehydrogenase 3 (NAD+) beta
	TTGTAAATGCG	6,00			kinesin family member 5B
-	ACCAGGCAAGG	2,00			hypothetical protein FLJ10783
	AGTGCCCCTCT	2,00			KIAA0552 gene product
	TGTGGCAAAGT	2,00		Hs.7243	ubiquitin specific protease 24
	TACCAGGAACC	2,00			NPD002 protein
	ATTITITCAAG	2,00		Hs.3833	3-prime-phosphoadenosine 5-prime-
		_,55	٥,ـ٥		phosphosulfate synt
3720	AGGCGCTTAGA	2,00	0.28	Hs.276	ESTs
	GGGATGGAAGG	2,00			hypothetical protein FLJ20185
	TCAAGGCCCCC	2,00			hypothetical protein DKFZp547M136
[_ <sup></sup>		-,	-,_5		similar to widely-
3723	AGACGCACTCT	2,00	0,28	Hs.139929	
	CTGAACTGTGA	2,00		Hs.121031	
	TCTGTAAAAAA	1,00			ESTs
-	AAGTCATCTAT	1,00			Homo sapiens mRNA; cDNA
لـــا					DKFZp564E153 (from clone DKF
<del></del>	TATTTTGAATA	1,00			translocation protein 1
3728	GGATCAAGTCC	1,00			damage-specific DNA binding protein 2
					(48kD)
3729	AAACTATGCAC	1,00	0,14		synuclein, alpha (non A4 component of
	110707				amyloid precur
	AAGTCCTGCAC	1,00		Hs.75871	protein kinase C binding protein 1
	ACGGTCCAGGA	1,00			cytidine deaminase
	TTTTAAAAAAA	1,00			ESTs
3/33	GGACACTCCTT	1,00	0,14	Hs.69360	kinesin-like 6 (mitotic centromere-
270	CATTOOCCATT	-		11a 04 4=5	associated kinesi
5/34	GATTGCGGATA	1,00	0,14		ESTs, Weakly similar to unknown
2705	CTCTTCCCC				[S.cerevisiae]
	GTGTTGGGGTG	1,00			anaphase-promoting complex subunit 7
-	TTGCTAAAGGC	1,00		Hs.5216	HSPC028 protein
	TATACAGATTG	1,00		Hs.4996	DKFZP564D166 protein
	CTGCAGAAAAA	1,00		Hs.4310	eukaryotic translation initiation factor 1A
	CAGTGTATTCG	1,00			CHMP1.5 protein
	CAGGGCTCACC	1,00		Hs.40411	ESTs
	GTGGGTCAGCT	1,00			ESTs
3/42	TAATGAATGAG	1,00	0,14	Hs.3164	nucleobindin 2

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2742	AATTACTTCCT	1.00	0.14	No 205440	CCT
	AATTACTTCCT	1,00		Hs.285449	
	GAGGGGTTTGC	1,00		Hs.285112	
	CCCTCCATTTG	1,00			truncated calcium binding protein
	CCCTTCAAAAA	1,00			Tubulin, alpha, brain-specific
	CTGGTTTAAAT	1,00			KIAA1228 protein
	TTTGTTTTAAG	1,00			eukaryotic translation initiation factor 4 gamma, 3
3749	GACAATGTATG	1,00	0,14	Hs.23767	guanine nucleotide binding protein (G protein), gamm
3750	CCACTGCCCTT	1,00	0,14	Hs.199695	hypothetical protein
3751	ATCCTCTGCGT	1,00			calcium/calmodulin-dependent protein kinase I
3752	GGAAATTGTCT	1,00	0,14	Hs.181112	HSPC126 protein
3753	CATTCACCATA	1,00	0,14	Hs.16552	Homo sapiens clone TCBA00758 mRNA sequence
3754	TTCAACAGGAA	1,00	0.14	Hs.158195	heat shock transcription factor 2
	ACTTCTGGAAC	1,00			KIAA0171 gene product
	GAAAATGCATC	1,00	0.14	Hs 154919	KIAA0625 protein
	TGAATAAAATG	1,00			matrix metalloproteinase 19
	GAGAAACTAGA	1,00	0.14	He 139120	ribonuclease P (30kD)
	GAGTGCAGGAC	1,00	0,14	He 120836	KIAA1028 protein
	GATTGGACTTG	1,00			galactokinase 2
	TGCCCTGAGAG	1,00	0,17	Ho 11127	PET112 (yeast homolog)-like
	CTTTTCTATGT			Hs.10711	
	TGAAGTGTATA	1,00 1,00			O-linked N-acetylglucosamine (GlcNAc) transferase (U
3764	AGACCAAAGTG	10,00	0.63	Hs 82646	heat shock 40kD protein 1
	AAACTCGAGCA	5,00			hypothetical protein FLJ20159
	TTACAGTCTTA	4,00			Homo sapiens mRNA; cDNA
2767	AGCTTGCGCTC	4.00	0.42	Un 161551	DKFZp434C0814 (from clone DK
	TTAAACTCTAA	4,00 3,00	0,42	Hs.226213	hypothetical protein FLJ20159 cytochrome P450, 51 (lanosterol 14-
0700	00400777000	0.00	- 0.00		alpha-demethylase
	CCACCTTTCCC	3,00			ESTs
	TAATAAAGGTG	76,00			ribosomal protein S8
	GTGGGTTGGCT	7,00	0,54	Hs.195432	aldehyde dehydrogenase 2, mitochondrial
	CAGTTACAAAG	4,00		Hs.77508	glutamate dehydrogenase 1
3773	ATGTGAAGAGT	59,00	1,07	Hs.111779	secreted protein, acidic, cysteine-rich (osteonectin
3774	TGTATGCCGTC	2,00	0,27	Hs.83469	nuclear factor (erythroid-derived 2)-like 1
	TATGAATGCTG	2,00		Hs.81800	chondroitin sulfate proteoglycan 2 (versican)
3776	ATATGAATGTG	2,00	0.27	Hs.7862	hypothetical protein FLJ20312
	CACTTCCTCCT	2,00		Hs.6375	uncharacterized hypothalamus protein HT010
3778	TTGTTATATTG	2,00	0.27	Hs.5862	hypothetical protein
	GATGTCTTGTT	2,00		Hs.5637	ESTs
	GAGGAGGAGGT				
3/80	GAGGAGGAGGI	2,00	U,2/	Hs.29956	KIAA0460 protein

2701	GAGCCAACAAT	2,00	0.27	Un 283690	hypothetical protein
	TGAGATTTCTT	2,00			CGI-150 protein
		2,00			step II splicing factor SLU7
	GAGCTGCATCA	5,00			melanoma antigen, family D, 1
	GTGCTCTGTAC				
	TTCTTGTGGCG	45,00			ribosomal protein S11
3/86	GTGATGTACGG	3,00	0,36	Hs.6639	Homo sapiens cDNA FLJ20818 fis, clone ADSE00627
3787	AAAGCATTTTC	3,00	0,36	Hs.6406	MO25 protein
3788	CGGCTGCCCAC	3,00	0,36	Hs.63236	synuclein, gamma (breast cancer- specific protein 1)
3789	ACCAGGCCACC	3,00	0,36	Hs.12068	carnitine acetyltransferase
	TTGCCCAGCAC	7,00		Hs.23954	cerebral cell adhesion molecule
3791	TGCTGAATCAG	8,00	0,57	Hs.2853	poly(rC)-binding protein 1
-	GCCTTGATCTC	3,00			DKFZP586E0820 protein
	ACAAAGGGCCC	3,00		Hs.7416	KIAA0397 gene product
	CCTCTGCACTC	3,00		Hs.265124	
	AGCCACTGCGC	9,00	0,59		EST, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
3796	ATTATCCAGGG	9,00	0.50	He 182225	RNA binding motif protein 3
	TACCAAGACCC	6,00		Hs.3059	coatomer protein complex, subunit beta
	GGCCGCGTTCG	36,00		Hs.5174	ribosomal protein S17
	ATCCTCCCTAT				RAP1A, member of RAS oncogene
		2,00			family
	GCAGCTCAAAG	2,00			alpha-2-macroglobulin
-	CTCAAGCGGCT	2,00			HIV-1 Tat interactive protein, 60 kDa
3802	GTCCGGAGTCT	2,00			protein phosphatase methylesterase-1
3803	AGCCACCACAG	2,00	0,27	Hs.245474	ESTs, Moderately similar to ALU5_HUMAN ALU SUBFAMILY
3804	AGCCAATTAAA	2,00	0,27	Hs.207625	
	GTGTAGTTGAG	2,00			adenylate kinase 2
	ACCCCAGCAAC	2,00			Homo sapiens cDNA FLJ20263 fis, clone COLF7804, high
3807	GCGATTAATTA	2,00	0,27	Hs.149436	kinesin family member 5B
3808	TCAAATGCAAA	2,00	0,27	Hs.116875	KIAA0156 gene product
3809	GGCAGAGACCC	1,00			hypothetical protein
3810	CTTCAAAAAAA	1,00	0,14	Hs.9667	butyrobetaine (gamma), 2-oxoglutarate dioxygenase (g
3811	CCGACGTCTCC	1,00	0,14	Hs.94943	ESTs
	CTGTGTTTGTT	1,00		Hs.9460	ESTs
	GGAGCTTAGAA	1,00		Hs.9194	putative glialblastoma cell differentiation- related
3814	GAAGATCCAGC	1,00	0,14	Hs.81875	growth factor receptor-bound protein 10
	AGGAAAAAAAC	1,00		Hs.79306	eukaryotic translation initiation factor 4E
	AACCAAAGGAA	1,00		Hs.78524	TcD37 homolog
	GAGCCAAGTGA	1,00		Hs.76536	transducin (beta)-like 1
	TCAAATCTTTG	1,00		Hs.75344	ribosomal protein S4, X-linked
	GCTTCCAGCTT	1,00		Hs.7407	KIAA1065 protein
	AAGGACACATC	1,00		Hs.6217	ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB

3822   TTCTCTTTCAA	3821	AGTAGTATGAA	1,00	0.14	Hs.56066	fibroblast growth factor 2 (basic)
BKZP434N0317 (from clone DK   3823 CTGTTCATCTA   1,00   0,14   Hs.51043   hexosaminidase B (beta polypeptida   3824 CCTGGAACCCC   1,00   0,14   Hs.4076   CTD (carboxy-terminal domina, RN/polymerase II, pol   3825 GCTGGAGAGTT   1,00   0,14   Hs.3447   DKFZP564K1964 protein   3826 TITTAGACAGC   1,00   0,14   Hs.283779   hypothetical protein DKFZp434J101   3828   ATAGCCTCTTA   1,00   0,14   Hs.283779   hypothetical protein DKFZp434J101   3829 TTGGGATGGGA   1,00   0,14   Hs.279799   putative zinc finger protein NY-REN antigen   3830   AGGTCAGGGGA   1,00   0,14   Hs.27328   EST, Moderately similar to   ALU2   HUMAN ALU SUBFAMILY   3831 TTGGCTGGGCT   1,00   0,14   Hs.26333   JM1 protein   3832 TGGGGAGCCG   1,00   0,14   Hs.26333   JM1 protein   3834 GGGGGGGGGGTT   1,00   0,14   Hs.261643   ESTs   3834 GGGGGGGGGTT   1,00   0,14   Hs.261643   ESTs   3835 CATTITTATAT   1,00   0,14   Hs.236030   SWI/SNF related, matrix associated actin dependent   3836 GCTCACTGTAG   1,00   0,14   Hs.236030   SWI/SNF related, matrix associated actin dependent   3836 GCTCACTGTAG   1,00   0,14   Hs.236183   EST   SM33 TTAAACCCACC   1,00   0,14   Hs.261649   Hs.261649   Hs.261649   Hs.261649   Hs.261649   STs   SM33 TTAAACCCACC   1,00   0,14   Hs.261664   Insulin-like growth factor 2 (somaton   A)   SM33 TTAAACCCACC   1,00   0,14   Hs.261664   Insulin-like growth factor 2 (somaton   A)   SM33 TTCCCAAGGCC   1,00   0,14   Hs.16766   Insulin-like growth factor 2 (somaton   A)   SM33 TTCCCAAGGCC   1,00   0,14   Hs.16766   Insulin-like growth factor 2 (somaton   A)   SM33 TTCCCAAGGCC   1,00   0,14   Hs.16766   Insulin-like growth factor 2 (somaton   A)   Insulin-like growth factor 2 (somaton   A)   SM33 TTCCCAAGGCC   1,00   0,14   Hs.16766   Insulin-like growth factor 2 (somaton   A)   SM33 TTCCCAAGGCC   1,00   0,14   Hs.16766   Insulin-like growth factor 2 (somaton   A)   Insulin-like growth factor 2 (somaton						
3823   CTGTTCATCTA			',"	•,		
3824   CCTGGAACCCC	3823	CTGTTCATCTA	1,00	0,14	Hs.51043	
Dolymerase II, pol   Dolymerase II, pol   Section   Se	$\overline{}$					
3826   TTTTAGACAGC			''-	•		
3827   TGTTGGTAAGT	3825	GCTGGAGAGTT	1,00	0,14	Hs.3447	<del></del>
3828   ATAGCCTCTTA	3826	TTTTAGACAGC	1,00	0,14	Hs.31720	hephaestin
antigen   antigen   3829   TTGGGATGGGA   1,00   0,14   Hs.278568   H factor (complement)-like 1   3830   AGGTCAGGGGA   1,00   0,14   Hs.277328   EST, Moderately similar to ALU2   HUMAN ALU SUBFAMILY   3831   TTGGCTGGGCT   1,00   0,14   Hs.274511   Homo sapiens mRNA; cDNA DKFZp586E121 (from clone DKF 3832   TGGGGAGCTCG   1,00   0,14   Hs.263633   JMI protein   3833   TCATTITITTG   1,00   0,14   Hs.261643   ESTs   3834   GGGGGGGGGTT   1,00   0,14   Hs.251664   insulin-like growth factor 2 (somaton A)   3835   CATTITATAT   1,00   0,14   Hs.236030   SWI//SNF related, matrix associated actin dependent   3836   GCTCACTGTAG   1,00   0,14   Hs.235183   EST   3837   TTAAACCCACC   1,00   0,14   Hs.26799   HSPC039 protein   3839   TGCCAAGGCC   1,00   0,14   Hs.197366   smoothened (Drosophila) homolog   3839   AGGCTTTAGCT   1,00   0,14   Hs.193082   ESTs   3840   GTAGCGCACAC   1,00   0,14   Hs.197366   smoothened (Drosophila) homolog   3841   GCGACAGTAAT   1,00   0,14   Hs.169387   KIAA0539   gene product   3842   AAGTGTGTTTT   1,00   0,14   Hs.165986   testin   3842   CAGTAGAAGG   1,00   0,14   Hs.155829   KIAA0676   protein   3843   CAGTAGAAGG   1,00   0,14   Hs.155829   KIAA0676   protein   3845   GAGATTGAGGC   1,00   0,14   Hs.155829   KIAA0676   protein   3846   GAGATTGAGGC   1,00   0,14   Hs.155829   KIAA0939   protein   3847   CCCACCGGTGC   1,00   0,14   Hs.126557   ESTs   3848   CAGGAGCCCCT   1,00   0,14   Hs.126557   ESTs   3849   TAGCAATTGCA   1,00   0,14   Hs.126557   ESTs   3850   GATGCAGAGC   1,00   0,14   Hs.126557   ESTs   3851   TTGGCCTTTTA   1,00   0,14   Hs.10554   ESTs, Weakly similar to unknown   D.melanogaster   3853   TTGTCTGCTA   7,00   0,54   Hs.237225   ribosomal protein   S5 pseudogene   1   3855   AGGATGTGGGC   9,00   0,59   Hs.139648   KIAA0706   gene product   3856   CCCTTAGCTTT   11,00   0,64   Hs.233936   myosin, light polypeptide, regulatory   non-sarcomeri	3827	TGTTGGTAAGT	1,00			
3829   TTGGGATGGGA   1,00	3828	ATAGCCTCTTA	1,00	0,14	Hs.279799	
3830   AGGTCAGGGGA   1,00   0,14   Hs.277328   EST, Moderately similar to ALU2   HUMAN ALU SUBFAMILY						
ALU2_HUMAN ALU SUBFAMILY   3831   TTGGCTGGGCT   1,00   0,14   Hs.274511   Homo sapiens mRNA; cDNA DKFZp586E121 (from clone DKF 3832   TGGGGAGCTCG   1,00   0,14   Hs.26133   JM1 protein   3833   TCATTTTTTTG   1,00   0,14   Hs.261643   ESTs   3834   GGGGGGGGGTT   1,00   0,14   Hs.251664   insulin-like growth factor 2 (somaton A)   3835   CATTTTATAT   1,00   0,14   Hs.236030   SWI/SNF related, matrix associated actin dependent   3836   GCTCACTGTAG   1,00   0,14   Hs.235183   EST   3837   TTAAACCCACC   1,00   0,14   Hs.235183   EST   3838   TTCCCAAGGCC   1,00   0,14   Hs.197366   smoothened (Drosophila) homolog   3839   AGGCTTTAGCT   1,00   0,14   Hs.18759   KIAA0539 gene product   3841   GCGACAGTAAT   1,00   0,14   Hs.170853   ESTs   3842   AAGTGTGTTTT   1,00   0,14   Hs.169387   KIAA0036 gene product   3843   CCAGTAGAAGG   1,00   0,14   Hs.165986   testin   3844   TTTATTGCACA   1,00   0,14   Hs.155372   ESTs   3845   GAGATTGAGGC   1,00   0,14   Hs.155372   ESTs   3846   GAGCACTTCCT   1,00   0,14   Hs.15144   hypothetical protein DKFZp564O04: 3847   CCCACCGGTGC   1,00   0,14   Hs.15144   hypothetical protein DKFZp564O04: 3849   TAGCAATTGCA   1,00   0,14   Hs.128557   ESTs   3850   GATGCAGCACC   1,00   0,14   Hs.128557   ESTs   3851   TTGGCCTTTTA   1,00   0,14   Hs.105540   ESTs, Weakly similar to unknown   D.melanogaster]   3853   TTGTTCTGCTA   7,00   0,54   Hs.237225   ribosomal protein S5 pseudogene 1   3854   TTGGTCCTCTG   241,00   1,46   Hs.108240   Ribosomal protein S5 pseudogene 1   3854   TGGTCCTCTG   241,00   1,46   Hs.233936   KIAA0706   gene product   3856   CCCTTAGCTTT   1,00   0,54   Hs.233936   KIAA0706   gene product   None product   3856   CCCTTAGCTTT   1,00   0,54   Hs.233936   KIAA0706   gene product   3856   CCCTTAGCTTT   1,00   0,64   Hs.233936   KIAA0706   gene product   3856   CCCTTAGCTTT   1,00   0,64   Hs.233936   KIAA0706   gene product   3856   CCCTTAGCTTT   1,00   0,64   Hs.233936   KIAA0706   gene product   None product   3856   CCCTTAGCTTT   1,00   0,64   Hs.233936   KIAA0706   gene p						
DKFZp586E121 (from clone DKF   3832 TGGGGAGCTCG   1,00   0,14 Hs.26333	3830	AGGTCAGGGGA	1,00	0,14	Hs.277328	
3832   TGGGGAGCTCG	3831	TTGGCTGGGCT	1,00	0,14		
3833   TCATTTTTTG						
3834   GGGGGGGGTT						
A)						
actin dependent   3836 GCTCACTGTAG   1,00   0,14   Hs.235183 EST   3837 TTAAACCCACC   1,00   0,14   Hs.226799   HSPC039 protein   3838 TTCCCAAGGCC   1,00   0,14   Hs.197366   smoothened (Drosophila)   homolog   3839   AGGCTTTAGCT   1,00   0,14   Hs.193082   ESTs   3840   GTAGCGCACAC   1,00   0,14   Hs.18759   KIAA0539 gene product   3841   GCGACAGTAAT   1,00   0,14   Hs.170853   ESTs   3842   AAGTGTGTTT   1,00   0,14   Hs.169387   KIAA0036 gene product   3843   CCAGTAGAAGG   1,00   0,14   Hs.165986   testin   3844   TTATTGCACA   1,00   0,14   Hs.155829   KIAA0676   protein   3845   GAGATTGAGGC   1,00   0,14   Hs.152372   ESTs   3846   GAGCACTTCCT   1,00   0,14   Hs.15144   hypothetical protein DKFZp564O043   3847   CCCACCGGTGC   1,00   0,14   Hs.134901   ESTs   3848   CAGGAGCCCCT   1,00   0,14   Hs.12785   KIAA0939   protein   3849   TAGCAATTGCA   1,00   0,14   Hs.126557   ESTs   3850   GATGCAGCAGC   1,00   0,14   Hs.105280   ESTs, Weakly similar to unknown   [D.melanogaster]   3853   TTGTCTGCTA   7,00   0,54   Hs.237225   ribosomal protein   S5   pseudogene   1 3854   TTGGTCCTCTG   241,00   1,46   Hs.108124   ribosomal protein   L41   3855   AGGATGTGGGC   9,00   0,59   Hs.139648   KIAA0706   gene product   3856   CCCTTAGCTTT   11,00   0,64   Hs.233936   myosin, light polypeptide, regulatory   non-sarcomeri	3834	GGGGGGGTT	1,00			A)
3836   GCTCACTGTAG	3835	CATTITTATAT	1,00	0,14		
3837   TTAAACCCACC						
3838   TTCCCAAGGCC						
3839   AGGCTTTAGCT   1,00   0,14   Hs.193082   ESTs	-					
3840 GTAGCGCACAC						
3841   GCGACAGTAAT   1,00   0,14   Hs.170853   ESTs     3842   AAGTGTGTTTT   1,00   0,14   Hs.169387   KIAA0036 gene product     3843   CCAGTAGAAGG   1,00   0,14   Hs.165986   testin     3844   TTTATTGCACA   1,00   0,14   Hs.155829   KIAA0676 protein     3845   GAGATTGAGGC   1,00   0,14   Hs.152372   ESTs     3846   GAGCACTTCCT   1,00   0,14   Hs.15144   hypothetical protein DKFZp564O043     3847   CCCACCGGTGC   1,00   0,14   Hs.134901   ESTs     3848   CAGGAGCCCCT   1,00   0,14   Hs.12785   KIAA0939 protein     3849   TAGCAATTGCA   1,00   0,14   Hs.126557   ESTs     3850   GATGCAGCAGC   1,00   0,14   Hs.10554   ESTs, Weakly similar to unknown						
3842   AAGTGTGTTTT						
3843   CCAGTAGAAGG						
3844 TTTATTGCACA         1,00         0,14 Hs.155829 KIAA0676 protein           3845 GAGATTGAGGC         1,00         0,14 Hs.152372 ESTs           3846 GAGCACTTCCT         1,00         0,14 Hs.15144 hypothetical protein DKFZp564O043           3847 CCCACCGGTGC         1,00         0,14 Hs.134901 ESTs           3848 CAGGAGCCCCT         1,00         0,14 Hs.12785 KIAA0939 protein           3849 TAGCAATTGCA         1,00         0,14 Hs.126557 ESTs           3850 GATGCAGCAGC         1,00         0,14 Hs.117582 CGI-43 protein           3851 TTGGCCTTTTA         1,00         0,14 Hs.10554 ESTs, Weakly similar to unknown [D.melanogaster]           3852 GGGTCTGCTGT         1,00         0,14 Hs.105280 ESTs, Weakly similar to dJ963K23.2 [H.sapiens]           3853 TTGTTCTGCTA         7,00         0,54 Hs.237225 ribosomal protein S5 pseudogene 1           3854 TTGGTCCTCTG         241,00         1,46 Hs.108124 ribosomal protein L41           3855 AGGATGTGGGC         9,00         0,59 Hs.139648 KIAA0706 gene product           3856 CCCTTAGCTTT         11,00         0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri				0,14	Hs.169387	KIAA0036 gene product
3845 GAGATTGAGGC	-					
3846         GAGCACTTCCT         1,00         0,14         Hs.15144         hypothetical protein DKFZp564O043           3847         CCCACCGGTGC         1,00         0,14         Hs.134901         ESTs           3848         CAGGAGCCCCT         1,00         0,14         Hs.12785         KIAA0939 protein           3849         TAGCAATTGCA         1,00         0,14         Hs.126557         ESTs           3850         GATGCAGCAGC         1,00         0,14         Hs.117582         CGI-43 protein           3851         TTGGCCTTTTA         1,00         0,14         Hs.10554         ESTs, Weakly similar to unknown [D.melanogaster]           3852         GGGTCTGCTGT         1,00         0,14         Hs.105280         ESTs, Weakly similar to dJ963K23.2 [H.sapiens]           3853         TTGTTCTGCTA         7,00         0,54         Hs.237225 ribosomal protein S5 pseudogene 1           3854         TTGGTCCTCTG         241,00         1,46         Hs.108124 ribosomal protein L41           3855         AGGATGTGGGC         9,00         0,59         Hs.139648 KIAA0706 gene product           3856         CCCTTAGCTTT         11,00         0,64         Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
3847 CCCACCGGTGC         1,00         0,14 Hs.134901 ESTs           3848 CAGGAGCCCCT         1,00         0,14 Hs.12785 KIAA0939 protein           3849 TAGCAATTGCA         1,00         0,14 Hs.126557 ESTs           3850 GATGCAGCAGC         1,00         0,14 Hs.117582 CGI-43 protein           3851 TTGGCCTTTTA         1,00         0,14 Hs.10554 ESTs, Weakly similar to unknown [D.melanogaster]           3852 GGGTCTGCTGT         1,00         0,14 Hs.105280 ESTs, Weakly similar to dJ963K23.2 [H.sapiens]           3853 TTGTTCTGCTA         7,00         0,54 Hs.237225 ribosomal protein S5 pseudogene 1           3854 TTGGTCCTCTG         241,00         1,46 Hs.108124 ribosomal protein L41           3855 AGGATGTGGGC         9,00         0,59 Hs.139648 KIAA0706 gene product           3856 CCCTTAGCTTT         11,00         0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
3848 CAGGAGCCCCT         1,00         0,14 Hs.12785         KIAA0939 protein           3849 TAGCAATTGCA         1,00         0,14 Hs.126557 ESTs           3850 GATGCAGCAGC         1,00         0,14 Hs.117582 CGI-43 protein           3851 TTGGCCTTTTA         1,00         0,14 Hs.10554 ESTs, Weakly similar to unknown [D.melanogaster]           3852 GGGTCTGCTGT         1,00         0,14 Hs.105280 ESTs, Weakly similar to dJ963K23.2 [H.sapiens]           3853 TTGTTCTGCTA         7,00         0,54 Hs.237225 ribosomal protein S5 pseudogene 1           3854 TTGGTCCTCTG         241,00         1,46 Hs.108124 ribosomal protein L41           3855 AGGATGTGGGC         9,00         0,59 Hs.139648 KIAA0706 gene product           3856 CCCTTAGCTTT         11,00         0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
3849   TAGCAATTGCA						
3850 GATGCAGCAGC         1,00         0,14 Hs.117582 CGI-43 protein           3851 TTGGCCTTTTA         1,00         0,14 Hs.10554 ESTs, Weakly similar to unknown [D.melanogaster]           3852 GGGTCTGCTGT         1,00         0,14 Hs.105280 ESTs, Weakly similar to dJ963K23.2 [H.sapiens]           3853 TTGTTCTGCTA         7,00         0,54 Hs.237225 ribosomal protein S5 pseudogene 1           3854 TTGGTCCTCTG         241,00         1,46 Hs.108124 ribosomal protein L41           3855 AGGATGTGGGC         9,00         0,59 Hs.139648 KIAA0706 gene product           3856 CCCTTAGCTTT         11,00         0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
3851 TTGGCCTTTA 1,00 0,14 Hs.10554 ESTs, Weakly similar to unknown [D.melanogaster] 3852 GGGTCTGCTGT 1,00 0,14 Hs.105280 ESTs, Weakly similar to dJ963K23.2 [H.sapiens] 3853 TTGTTCTGCTA 7,00 0,54 Hs.237225 ribosomal protein S5 pseudogene 1 3854 TTGGTCCTCTG 241,00 1,46 Hs.108124 ribosomal protein L41 3855 AGGATGTGGGC 9,00 0,59 Hs.139648 KIAA0706 gene product 3856 CCCTTAGCTTT 11,00 0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
[D.melanogaster]  3852 GGGTCTGCTGT 1,00 0,14 Hs.105280 ESTs, Weakly similar to dJ963K23.2 [H.sapiens]  3853 TTGTTCTGCTA 7,00 0,54 Hs.237225 ribosomal protein S5 pseudogene 1  3854 TTGGTCCTCTG 241,00 1,46 Hs.108124 ribosomal protein L41  3855 AGGATGTGGGC 9,00 0,59 Hs.139648 KIAA0706 gene product  3856 CCCTTAGCTTT 11,00 0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri			1,00			
3852   GGGTCTGCTGT	3851	TTGGCCTTTTA	1,00	0,14		
3853 TTGTTCTGCTA   7,00   0,54 Hs.237225 ribosomal protein S5 pseudogene 1   3854 TTGGTCCTCTG   241,00   1,46 Hs.108124 ribosomal protein L41   3855 AGGATGTGGGC   9,00   0,59 Hs.139648 KIAA0706 gene product   3856 CCCTTAGCTTT   11,00   0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri	3852	GGGTCTGCTGT	1,00	0,14		ESTs, Weakly similar to dJ963K23.2
3854   TTGGTCCTCTG   241,00   1,46   Hs.108124   ribosomal protein L41   3855   AGGATGTGGGC   9,00   0,59   Hs.139648   KIAA0706 gene product   3856   CCCTTAGCTTT   11,00   0,64   Hs.233936   myosin, light polypeptide, regulatory non-sarcomeri	3853	TTGTTCTGCTA	7.00	0.54	Hs.237225	
3855 AGGATGTGGC 9,00 0,59 Hs.139648 KIAA0706 gene product 3856 CCCTTAGCTTT 11,00 0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
3856 CCCTTAGCTTT 11,00 0,64 Hs.233936 myosin, light polypeptide, regulatory non-sarcomeri						
					Hs.233936	myosin, light polypeptide, regulatory,
TANALLE LANGUAGE LA 2001 LA 2001 LA 2011 LA 2011 LA LA CALLELLA CALLEL ILLEHIDISHE DICHENI	3857	TTTCAGTGGGT	3,00	0.36		
3858 GCCCAGCCTG 3,00 0,36 Hs.180903 hypothetical protein						
3859 GTTTGGAGCTG 3,00 0,36 Hs.180533 mitogen-activated protein kinase kin						

	· -	<del></del>			Ia
0000	A A O T A A A A A A	47.00	0.07	11 55004	3
	AACTAAAAAAA	47,00		Hs.55921	glutamyl-prolyl-tRNA synthetase
	GACACCTCCTG	7,00		Hs.6455	RuvB (E coli homolog)-like 2
	CCTCCCTGATG	5,00		Hs.167013	
	AGGAATGGTAG	2,00		Hs.7911	KIAA0323 protein
	TGGTCCAGCGC	2,00		Hs.75627	CD14 antigen
	TGACTTATTAA	2,00		Hs.74649	cytochrome c oxidase subunit VIc
	TGTGTGTGACA	2,00			ESTs
	CCAAAGAGTAT	2,00			HSPC158 protein
3868	TTTCTTAATGT	2,00	0,27	Hs.233650	ESTs, Weakly similar to H beta 58
2222					homolog [H.sapiens
3869	TAAAACAAGAA	2,00	0,27	Hs.1369	decay accelerating factor for complement
		22.22			(CD55, Crom
	GACAAAAAAA	20,00		Hs.2953	ribosomal protein S15a
	AGGTCAAGAGA	5,00			KIAA1185 protein
	TCAATAAAGAA	8,00			glutaminyl-tRNA synthetase
	TGCTGGTGTGG	3,00			KIAA0864 protein
3874	CCAATTTACAA	3,00	0,36	Hs.274472	high-mobility group (nonhistone
	0.707.0				chromosomal) protein
3875	CTGTAGAAATG	3,00	0,36	Hs.215595	guanine nucleotide binding protein (G
0070					protein), beta
3876	AATGAATAAAA	3,00	0,36	Hs.118797	ubiquitin-conjugating enzyme E2D 3
2077	00040707000		0.00	11 40000	(homologous to ye
	GCGAGTCTCCG	3,00	0,36	Hs.10632	hypothetical protein DKFZp762M136
3878	GGGAAGTCACC	4,00	0,42	Hs.264428	tissue specific transplantation antigen
2070	CTCCCACCTCC	4 00	0.40	11- 4044	P35B
	GTGGCACCTGC	4,00		Hs.1244	CD9 antigen (p24)
3000	CTGACTTGTGT	9,00	0,59	Hs.77961	major histocompatibility complex, class I, B
3881	AGGCTACGGAA	135,00	1.21	Hs.119122	ribosomal protein L13a
	AAGAGGTTTGC	6,00			transmembrane protein (63kD),
		-,	-,		endoplasmic reticulum/
3883	TGGCTAGTGTT	11,00	0.63		proteasome (prosome, macropain)
		·	·		subunit, beta type,
3884	TTTTGTTTTGT	2,00	0,27		transmembrane 4 superfamily member
					(tetraspan NET-7)
3885	GGGTAATGTGA	2,00	0,27	Hs.76907	HSPC002 protein
	ATCCGGACCCT	2,00			growth arrest and DNA-damage-
					inducible 34
3887	TGCCTGGAACT	2,00	0,27	Hs.6820	ESTs, Weakly similar to putative
					[C.elegans]
3888	GGCTGCAGTCT	2,00	0,27	Hs.48320	DKFZP566B1346 protein
3889	CTCTGATAACT	2,00			zinc metalloproteinase, STE24 (yeast,
					homolog)
3890	GTGTCTCCCGC	2,00	0,27	Hs.219614	f-box and leucine-rich repeat protein 11
	GCGAACCCCCC	2,00		Hs.211862	
	TTTTCTTAAA	2,00		Hs.197955	KIAA0704 protein
3893	AAGAATCAAAA	2,00			NADH dehydrogenase (ubiquinone) 1
L					beta subcomplex, 1

3894	GTGCTGATTCT	2,00	0,27	Hs.1640	collagen, type VII, alpha 1 (epidermolysis
3895	AACCAATCTGG	2,00	0,27		bullosa, cytochrome P450, subfamily I (dioxin-
L					inducible), pol
3896	AGGGAAAAAA	5,00	0,46	Hs.215595	guanine nucleotide binding protein (G protein), beta
3897	AATTTGCAACA	6,00	0,50	Hs.75258	H2A histone family, member Y
3898	TCAAATGTCAG	3,00	0,36	Hs.89474	ADP-ribosylation factor 6
3899	ATTTTGTGCAA	3,00	0,36	Hs.8750	uncharacterized bone marrow protein BM045
3900	ттсттетттте	8,00	0,56	Hs.74621	prion protein (p27-30) (Creutzfeld-Jakob disease, Ge
3901	GCCACTACCCC	4,00	0,42	Hs.71475	hypothetical protein
3902	GAGCAAACGGA	4,00	0,42	Hs.108847	Homo sapiens chromosome 19, cosmid R26445
3903	TCATACAGTTT	1,00	0,14	Hs.94986	ribonuclease P (38kD)
	GAGGGAAAAAG	1,00		Hs.9082	nucleoporin p54
-	CTGATGAATTC	1,00	0,14	Hs.86386	myeloid cell leukemia sequence 1 (BCL2-related)
3906	AAGCGCTACCT	1,00	0,14	Hs.83848	triosephosphate isomerase 1
3907	GAAGGCTTTAT	1,00	0,14	Hs.8182	KIAA0796 protein
3908	ACTTGATTCAA	1,00	0,14	Hs.80905	Ras association (RalGDS/AF-6) domain family 2
3909	CATTGTAATTC	1,00	0,14		KIAA0164 gene product
3910	ATTTTGTAACC	1,00			retinoblastoma-like 2 (p130)
3911	CATTCTCCTAG	1,00	0,14		SKI-INTERACTING PROTEIN
	TCCTGCTCATT	1,00		Hs.7844	golgi autoantigen, golgin subfamily b, macrogolgin (
3913	TGGCCTTTTTG	1,00	0,14	Hs.75761	SFRS protein kinase 1
	TTTTACATCTT	1,00			thyroid hormone receptor interactor 10
3915	CTGTGTAAAGG	1,00		Hs.7314	KIAA0614 protein
	CAGGGAATGCC	1,00		Hs.58598	KIAA1266 protein
3917	GCAGAGCAGTC	1,00			lymphoblastic leukemia derived sequence 1
3918	TTCACTAATTG	1,00			Homo sapiens mRNA; cDNA DKFZp434O0227 (from clone DK
3919	CTATTCCATTT	1,00	0,14	Hs.43505	inhibitor of kappa light polypeptide gene enhancer i
3920	TTTGTATGGGA	1,00	0,14	Hs.431	murine leukemia viral (bmi-1) oncogene homolog
3921	ATGTGGGTCTA	1,00	0,14	Hs.42392	ESTs
3922	GAAACTAGATC	1,00	0,14	Hs.37883	ESTs
3923	TTGTAACGTGT	1,00	0,14	Hs.3610	KIAA0205 gene product
3924	CTATGCATCAG	1,00	0,14	Hs.30670	hepatocellular carcinoma-associated antigen 66
3925	TTTGAGTTCTG	1,00	0,14	Hs.29494	PRO1912 protein
	CTACTGCAGTC	1,00			peroxisomal trans 2-enoyl CoA reductase; putative sh
3927	CTCCTCAGGGC	1,00	0.14	Hs.28088	SGC32445 protein

0000	0107070000	4 001	0.44		11710 0 4 4 4 6
	GACTCTGGCCC	1,00			KIAA1442 protein
3929	TAATGGGAGTC	1,00	0,14	Hs.264330	N-acylsphingosine amidohydrolase (acid
					ceramidase)-l
	CCTGTGATCTC	1,00		Hs.254176	
	ATTAGCAGAGT	1,00	<u>0,14</u>	Hs.249982	cathepsin B
3932	GTGGTACATAC	1,00	0,14	Hs.232120	ESTs, Weakly similar to ALU1_HUMAN
					ALU SUBFAMILY J S
3933	CCCATCAATAA	1,00	0,14	Hs.220689	Ras-GTPase-activating protein SH3-
					domain-binding pro
3934	CTATCAGTCTC	1,00	0,14	Hs.211601	mitogen-activated protein kinase kinase
					kinase 12
3935	TGTTCCAAGGC	1,00	0,14	Hs.211569	G protein-coupled receptor kinase 5
3936	ATTAATGAATC	1,00	0,14	Hs.18259	putative ATP(GTP)-binding protein
3937	AGAGGTTGATG	1,00	0,14	Hs.181244	major histocompatibility complex, class I,
<u> </u>					A The state of the
3938	TCAACATCTAG	1,00	0,14	Hs.171734	protein phosphatase 2, regulatory
<b>}</b>					subunit B (B56), g
3939	AAACTAGTTGC	1,00	0,14		myosin phosphatase, target subunit 1
3940	CCAGCTCCTTG	1,00			hypothetical protein DKFZp762L0311
3941	ATTCTCTCAGG	1,00			hypothetical protein FLJ20159
	AGAACAGACCA	1,00			ARP1 (actin-related protein 1, yeast)
		.,	-,		homolog A (cen
3943	TAGTCTGGAGT	1,00	0.14	Hs.152981	CDP-diacylglycerol synthase
		. ","	-,		(phosphatidate cytidylyl
3944	TGAAAGTCCTG	1,00	0.14	Hs. 152707	glioblastoma amplified sequence
	CTACTGAAAAA	1,00		Hs.15248	
	AATGTTTGTGA	1,00		Hs.135835	
	TTTGATTCTGT	1,00		Hs.12245	
$\overline{}$	TCTCTAAGCCA	1,00			3-hydroxy-3-methylglutaryl-Coenzyme A
00.0	1010171100071	1,00	0, 17		reductase
3949	GGAGCTTGAGG	1,00	0.14		Notch (Drosophila) homolog 4
	TGAGTGAATTC	1,00			Homo sapiens mRNA; cDNA
0000	IOAOIOAAIIO	1,00	0, 14		DKFZp586O2124 (from clone DK
3951	TGCTAATTGTA	4,00	0.41		Homo sapiens mRNA; cDNA
000	IOOIATIOIA	7,00	0,41	115.7 1900	DKFZp564F053 (from clone DKF
3952	TCCAGCCCCTG	5,00	0.46	Hs.24956	ESTs, Weakly similar to AF118023 1
0002	100/1000010	. 3,00	0,40		SH3 domain-bindin
3053	TATTTATTCCT	5,00	0.46		CGI-96 protein
	GAGAAGACTTC	2,00			prolyl endopeptidase
	AGTAAACCATC				Homo sapiens mRNA; cDNA
3933	AGIAAACCAIC	2,00	0,21		
2056	TCTTTCCAAAC	2.00	0.27		DKFZp586C1723 (from clone DK
2920	TCTTTGCAAAG	2,00	0,27		Homo sapiens mRNA; cDNA
2057	ACACTTCTTGG	2.00	0.27		DKFZp586C1817 (from clone DK
393/	ACACHOLIGG	2,00	0,27		ESTs, Moderately similar to EFGM_RAT
2050	<u> </u>	2.00	<del></del>		ELONGATION FACT
3938	AAAACAAAAA	2,00	0,27		solute carrier family 7 (cationic amino
2050	CATOTTOAGGA	40.00			acid transpo
_	CATCTTCACCA	19,00			ribosomal protein S25
3960	AGGCTGGATGC	4,00	0,41	Hs.5898	KIAA0668 protein

3961	TTGGCCAGGGT	4,00	0.41	Hs.209396	ESTs, Weakly similar to plakophilin 2b
	1,0000,10001	.,,55	•,		[H.sapiens]
3962	ATAGACGCAAT	11,00	0,63	Hs.6353	MORF-related gene 15
	TATCACTCTGT	5,00			male-enhanced antigen
	CAAGAGGCAAA	7,00		Hs.5734	KIAA0679 protein
	ATGTACTCTGG	8,00			IMP (inosine monophosphate)
		5,55	0,00		dehydrogenase 2
3966	GTGGTGCGCGC	3,00	0.35	Hs.252075	Homo sapiens mRNA; cDNA
			,,,,		DKFZp434D179 (from clone DKF
3967	TCCTGCCCCAT	22,00	0,78	Hs.171814	parathymosin
3968	TTGTTATTGCC	4,00			annexin A7
3969	GGGCCCCCTGG	2,00			glycophorin C (Gerbich blood group)
	TTATGTTTAAT	2,00			lumican
	AGTAGGAGGGA	2,00			hypothetical protein FLJ20550
	TTGTATCAGAA	2,00			FK506 binding protein 12-rapamycin
		·	·		associated protei
3973	GTGCAGTCCTC	2,00	0,27	Hs.19223	ESTs, Weakly similar to R26660_1,
					partial CDS [H.sap
3974	TTTTGCTACAG	2,00	0,27	Hs.171545	HIV-1 Rev binding protein
3975	TGAAACTTTTC	2,00	0,27	Hs.107528	androgen induced protein
3976	CTGGCCGCAAG	5,00	0,46	Hs.74649	cytochrome c oxidase subunit VIc
3977	GACTCTCTCAG	3,00	0,35	Hs.178576	similar to Bos taurus P14 protein
3978	AGCTGGGTTGG	3,00	0,35	Hs.131731	hypothetical protein FLJ11099
3979	TCTTCTAAAAA	3,00	0,35	Hs.108112	histone fold protein CHRAC17; DNA
					polymerase epsilon
3980	GTGGGGTCTCT	1,00	1,26		ESTs, Weakly similar to predicted using
					Genefinder [
3981	AATTTGAAAAA	1,00	1,26		ESTs, Highly similar to CMP-N-
					acetylneuraminic acid
	AGTGCATTGTA	1,00			ESTs
3983	CACTGTAGTCC	1,00	1,26	Hs.98952	Human DNA sequence from clone RP1-
2224	0071000100				39G22 on chromosom
	GGTAGGCAGGG	1,00			ESTs
	GTTAAATGACT	1,00			ESTs
3986	TAACTTTTTC	1,00	1,26	Hs.9667	butyrobetaine (gamma), 2-oxoglutarate
0007	00407040407	4.00	4 00		dioxygenase (g
	CCAGTGACACT	1,00		Hs.96023	CD19 antigen
3988	CAGATGCATCA	1,00	1,26	Hs.94695	ESTs, Moderately similar to AF238978_1 pIFI27-like p
3989	GGATGTCCTAT	1,00	1,26	Hs.94479	transmembrane protein 1
3990	TTGGGCAGGAA	1,00	1,26	Hs.92254	hypothetical protein FLJ20163
3991	TGGCAAAATGA	1,00	1,26	Hs.91728	polymyositis/scleroderma autoantigen 1 (75kD)
3992	TTTTAACTTTG	1,00	1,26	Hs.90753	Tat-interacting protein (30kD)
	TGTGAATATGC	1,00		Hs.89679	interleukin 2
	TGCAAGGCTTT	1,00		Hs.89418	prostaglandin F receptor (FP)
	TGGGAGCTCAG	1,00		Hs.88411	DNA segment on chromosome 6
		, ,	,_ •		(unique) 49 expressed se
3996	ATAAAGTCTAT	1,00	1,26	Hs.87745	ESTs, Weakly similar to Unknown

	Γ				[H.sapiens]
2007	GCCACTGAACC	1,00	1 26	Hs.87125	EH-domain containing 3
		1,00		Hs.857	retinol-binding protein 3, interstitial
	TATATACACAT	1,00		Hs.85112	insulin-like growth factor 1 (somatomedia
	GTGTGCATCTT				(C)
	GCACTCTATGT	1,00	<del></del>	Hs.846	interleukin 8 receptor, beta
4001	GGAAACCTTTT	1,00	1,26	Hs.8373	ESTs
4002	ATGTTAACATC	1,00	1,26	Hs.82719	Homo sapiens mRNA; cDNA DKFZp586F1822 (from clone DK
4003	TGTTTAAAAAT	1,00	1,26	Hs.82689	tumor rejection antigen (gp96) 1
4004	CATATTCACAT	1,00	1,26	Hs.82582	integrin, beta-like 1 (with EGF-like repeat domains)
4005	TGGAGACTTGC	1,00	1,26	Hs.82283	5-methyltetrahydrofolate-homocysteine methyltransfer
4006	TACCTAATTAT	1,00	1,26	Hs.82223	chordin-like
	CTTGTACCAGA	1,00		Hs.82124	laminin, beta 1
	TATAAGAAAAA	1,00		Hs.80306	Homo sapiens mRNA, clone:RES4-4
	CTACTGTACCC	1,00		Hs.80120	UDP-N-acetyl-alpha-D-
		.,,-	,		galactosamine:polypeptide N-ace
4010	ATCTACCTTGG	1,00	1,26	Hs.79093	EBNA-2 co-activator (100kD)
	GTGAAACTGTT	1,00		Hs.79090	exportin 1 (CRM1, yeast, homolog)
	GTAGAAAAAAG	1,00		Hs.78890	numb (Drosophila) homolog
	TTTGCACACAC	1,00		Hs.78518	natriuretic peptide receptor B/guanylate cyclase B (
4014	ACAATCTGTAG	1,00	1 26	Hs.77899	tropomyosin 1 (alpha)
	GTGCTATTTTG	1,00		Hs.77873	Homo sapiens mRNA full length insert cDNA clone EURO
1016	CTGAACGCCAT	1.00	1 26	Hs.77694	KIAA0429 gene product
	GAGGTGATGGT	1,00			major histocompatibility complex, class
		1,00		Hs.76807	II, DR alpha
	GGCACAGAGAG	1,00	1,26	Hş.76716	pre-alpha (globulin) inhibitor, H3 polypeptide
	TCCCTGAAAAG	1,00		Hs.76364	allograft inflammatory factor 1
4020	ATCTCAACTCA	1,00	1,26	Hs.75813	polycystic kidney disease 1 (autosomal dominant)
4021	GGTGTCTCCAG	1,00	1,26	Hs.75510	annexin A11
4022	GTGGGTGCTTT	1,00	1,26	Hs.75104	RNA-binding protein S1, serine-rich domain
4023	GAGCCAGAGCG	1,00	1.26	Hs.75080	LIM and SH3 protein 1
	AGGGGGCTGAA	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	GTGACCCCAAA	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	TTCAGCAGCAG	1,00		Hs.7239	SEC24 (S. cerevisiae) related gene
					family, member B
	TTTTGAACAGC	1,00		Hs.72249	protease-activated receptor 3
	GAACTTGAAGT	1,00		Hs.7165	zinc finger protein 259
4029	GTAAGCGCTGT	1,00	1,26 	Hs.70725	gamma-aminobutyric acid (GABA) A receptor, pi
4030	AGTTGACTCCC	1,00	1,26	Hs.69606	ESTs
4031	GCGTTTCTCCA	1,00	1,26	Hs.6853	carbohydrate (N-acetylglucosamine 6-O)

	<del>,                                    </del>			<del></del>	sulfotransfer
4032	TGAGCTTGGAA	1,00	1.26	Hs.6820	ESTs, Weakly similar to putative
	10,1001100,01	,,00	.,_0	1.0.0020	[C.elegans]
4033	TGTGGGAATAG	1,00	1 26	Hs.6783	ESTs
	AACTCACAGTT	1,00		Hs.66072	ESTs
	TGCAACATAAA	1,00		Hs.66020	ESTs
	ACTTGAAAAAA	1,00		Hs.64837	cystinosis, nephropathic
$\overline{}$	GGACTGCTGGG	1,00		Hs.6434	hypothetical protein DKFZp761F2014
	CCCAGCAGGTT			Hs.6351	
		1,00			cleavage and polyadenylation specific factor 4, 30kD
	CAAACTTTTGG	1,00		Hs.63348	DKFZP586M121 protein
4040	AAAAGATCCAG	1,00	1,26	Hs.6318	peroxisomal short-chain alcohol dehydrogenase
4041	ACATTCTACAA	1,00	1,26	Hs.61957	ESTs
4042	ATTICTIGITI	1,00	1,26	Hs.61508	Homo sapiens cDNA FLJ10991 fis, clone PLACE1002072
4043	CACATTTCTGT	1,00	1,26	Hs.6136	KIAA1391 protein
4044	TAATTGATCAT	1,00	1,26	Hs.606	ATPase, Cu++ transporting, alpha
					polypeptide (Menkes
4045	GCACACTTGCA	1,00	1,26	Hs.60440	ESTs, Weakly similar to serin protease with IGF-bind
4046	CGCTCTAGGCT	1,00	1 26	Hs.59970	ESTs
	GAGGAGTGCAT	1,00		Hs.59395	Homo sapiens clone IMAGE:112574
					mRNA sequence
	TTCTTGGACTT	1,00		Hs.58882	Microfibril-associated glycoprotein-2
_	AGCCTGACCCC	1,00		Hs.58611	ESTs
4050	CAGAGTCCCTG	1,00	1,26	Hs.57922	X-prolyl aminopeptidase (aminopeptidase P) 2, membra
4051	CTTCCTCTTGC	1,00	1,26	Hs.57836	ESTs
4052	TTGACGCTGTA	1,00	1,26	Hs.57697	hyaluronan synthase 1
4053	GTAATAAGTGT	1,00	1,26	Hs.5476	serine protease inhibitor, Kazal type, 5
4054	ATGGGCCCTAG	1,00		Hs.5378	spondin 1, (f-spondin) extracellular matrix protein
4055	CCCAATGGCCC	1,00	1,26	Hs.5181	proliferation-associated 2G4, 38kD
	ATAATGCCATT	1,00		Hs.49927	Homo sapiens mRNA; cDNA
			·		DKFZp434H1720 (from clone DK
4057	GCTTGACAAGT	1,00	1,26	Hs.49599	Homo sapiens mRNA; cDNA DKFZp434G0827 (from clone DK
4058	GCTTAACCGCC	1,00	1.26	Hs.48962	ESTs
	TATTTACTTTG	1,00		Hs.48778	Homo sapiens mRNA; cDNA
			1,20		DKFZp586O0221 (from clone DK
	ACCCAGGGGAG	1,00	1,26	Hs.46794	ESTs
	TAGAAAACTTC	1,00	1,26	Hs.44649	ESTs
4062	AGAGTGAAAAA	1,00	1,26	Hs.44592	beta-1,4 mannosyltransferase
4063	CCCTTTACTGC	1,00	1,26	Hs.44441	ESTs
	CTGACCTTTTA	1,00		Hs.44296	ESTs, Highly similar to dJ842G6.1 [H.sapiens]
4065	TCATCTTTGCC	1,00	1.26	Hs.43658	DKFZP586L151 protein
	GGCCCATCCCT	1,00		Hs.42853	cAMP responsive element binding

$\overline{}$					protein-like 1
4067	TGAAATACAAA	1,00	1 26	Hs.42745	ESTs
	AATCCTTACTC	1,00		Hs.41735	purinergic receptor P2X, ligand-gated ion channel, 1
4069	CTGGGCCATTG	1,00	1,26	Hs.4	alcohol dehydrogenase 2 (class I), beta polypeptide
4070	CTGTAATATTC	1,00	1,26	Hs.38703	ESTs
4071	CCACAGCCTAT	1,00	1,26	Hs.36131	collagen, type XIV, alpha 1 (undulin)
4072	GGTGAGATTGT	1,00	1,26	Hs.33983	ESTs, Weakly similar to ALU6_HUMAN ALU SUBFAMILY SP
4073	GAATCTCTCCC	1,00	1,26	Hs.32425	ESTs
4074	ATGGACCAAAG	1,00	1,26	Hs.32345	ESTs
	AATCTTGTTAA	1,00		Hs.32343	ESTs
	GCCTCTTTTCT	1,00		Hs.31439	serine protease inhibitor, Kunitz type, 2
	AATGGCCTTGA	1,00		Hs.31431	ESTs
	GGCATTGGTCA	1,00		Hs.3128	polymerase (RNA) II (DNA directed) polypeptide H
	GTCTTGTAATC	1,00			putative DNA binding protein
	TAGATAGCACA	1,00		Hs.31016	putative DNA binding protein
	TCTCAAAACAA	1,00		Hs.30591	ESTs
4082	AAGAACTCTGC	1,00	1,26		v-maf musculoaponeurotic fibrosarcoma (avian) oncoge
	AGTAATTTAAA	1,00			ESTs
4084	CATTCTTTCTT	1,00	1,26	Hs.30035	splicing factor, arginine/serine-rich (transformer 2
4085	GGCAGCAGGAC	1,00	1,26	Hs.29759	RNA POLYMERASE I AND TRANSCRIPT RELEASE FACTOR
4086	CTTGTTATTCA	1,00	1,26	Hs.29664	Human DNA sequence from clone 682J15 on chromosome 6
4087	CACTCTGTTTG	1,00			toll-like receptor 3
	TTCCCCCACCT	1,00	1,26	Hs.286217	KIAA0685 gene product
	CCTCAGCCTCT	1,00	1,26	Hs.285657	ESTs
	CTTTTTTCATT	1,00	1,26	Hs.285122	ESTs, Weakly similar to S53869 laminin beta-2 chain
	TCTCCCACCCC	1,00		Hs.285042	
	CACCTGTAAGC	1,00		Hs.284664	
	GCAGCGGGGAT	1,00		Hs.284422	
	GTIGTTTTIGG	1,00			complement component 3
	CCAGCCTCTGT	1,00			KIAA1535 protein
	TGTGGGAGCCA	1,00			chromosome 3 open reading frame 4
	CTTGTAGCCCT	1,00			hypothetical protein PRO3077
	TTCCAGAAGCA	1,00		Hs.283600	
	CATCCACCTGG	1,00		Hs.283527	
	AAAAGAATACT	1,00		Hs.282887	
	TATATCTCTCT	1,00		Hs.28273	
	TTGGCAGCAAT	1,00		Hs.280531	<del></del>
	GTAAAATTTGA	1,00			mitochondrial carrier homolog 1
	TATATAATGTG	1,00			zinc finger protein 222
4105	GAAACCCAGGG	1,00	1,26	Hs.279813	hypothetical protein

4106	CCACAGCTCTC	1,00	1 26	He 279671	katanin p60 (ATPase-containing) subunit
					A 1
	TTTGTTCCTGA	1,00		Hs.279531	
4108	GCAGAAGCACC	1,00	1,26	Hs.279396	EST, Weakly similar to AF119917_56 PRO2729 [H.sapien
4109	AGTGGTGAGGG	1,00	1,26	Hs.279298	ESTs
4110	TCAGCCCTGGC	1,00	1,26	Hs.278550	ESTs, Weakly similar to KIAA0940 protein [H.sapiens]
4111	CCTCTCATCCA	1,00	1,26	Hs.278028	ESTs
4112	GTGGTGCGCTT	1,00	1,26	Hs.277808	EST
4113	CACCCTGTAGT	1,00	1,26	Hs.277531	ESTs
	GTGAGACCTAA	1,00	1,26	Hs.277186	ESTs, Weakly similar to ALU6_HUMAN ALU SUBFAMILY SP
4115	ACTITGTAAAA	1,00	1,26	Hs.274976	EST
	TGCTGGGAACT	1,00	1,26	Hs.274667	EST
4117	CACCTATAGTA	1,00	1,26	Hs.274510	Homo sapiens mRNA; cDNA
					DKFZp564B032 (from clone DKF
4118	GCAGGTACGCT	1,00	1,26	Hs.274429	Homo sapiens cDNA FLJ11192 fis, clone PLACE1007618,
4119	CCAGGGGGGCC	1,00	1,26	Hs.272813	dual oxidase 1
4120	CTGGCCAACGT	1,00			hypothetical protein FLJ20306
4121	CCCAAACGCGC	1,00	1,26	Hs.272572	hemoglobin, alpha 2
4122	CCCAACGCGTT	1,00	1,26	Hs.272572	hemoglobin, alpha 2
	CCCTTTTAAAA	1,00	1.26	Hs.272325	Homo sapiens mRNA; cDNA
		, ,	.,		DKFZp434A2322 (from clone DK
4124	ATTGCTCCAAT	1,00	1,26	Hs.272146	ESTs, Moderately similar to ALU5_HUMAN ALU SUBFAMILY
4125	AAACAGTGTCT	1,00	1,26	Hs.270618	ESTs, Weakly similar to KIAA0822 protein [H.sapiens]
4126	AGCTACCACCC	1,00	1.26	Hs.269541	
	CCCCTGCACTG	1,00			ESTs, Moderately similar to Ibd1
		1,55	,,	110.20 1010	[H.sapiens]
4128	ATCCACAATGG	1,00	1,26	Hs.264428	tissue specific transplantation antigen P35B
4129	AAACACGGCAA	1,00	1,26	Hs.264221	ESTs
4130	TTGCCCAAAAA	1,00	1,26	Hs.26339	ESTs
4131	CCTGTAATTTT	1,00	1,26	Hs.259691	Homo sapiens mRNA; cDNA DKFZp761E0311 (from clone DK
4132	TGAGAAAACAG	1,00	1.26	Hs.259315	
	GGGCCCTGGGC	1,00			ESTs, Weakly similar to PI-3 kinase
					[H.sapiens]
	CCTGAAGCCTG	1,00			striatin, calmodulin-binding protein
	AATATGCATCC	1,00		Hs.258400	
	GATCAAAGAAG	1,00		Hs.257620	
	TATTTTGGTGC	1,00		Hs.257540	
	TGTTTTAGTTC	1,00	1,26	Hs.257441	EST
	AGTTGCTCTGG	1,00	1,26	Hs.257249	ESTs
4140	AGATTAACATT	1,00	1,26		ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S

4141	CATTCATTCAT	1,00	1,26	Hs.255906	ESTs
4142	GGTGTTCAGGG	1,00	1,26	Hs.255530	ESTs
4143	CACTCAAAATA	1,00	1,26	Hs.255374	ESTs
4144	GCCTGAGGCTT	1,00	1,26	Hs.255308	EST
4145	TAGCCGCTGGG	1,00	1,26	Hs.255176	ESTs, Moderately similar to
)		· )			CD45_HUMAN LEUKOCYTE COM
4146	TGCTTGGCCAG	1,00	1,26	Hs.255170	EST
	AATGGGAAGAT	1,00	1,26	Hs.255037	EST
	CAATGTAAAAG	1,00		Hs.254881	
	GCCCTAGAGCA	1,00			aquaporin 7
	CAACCATTTGC	1,00		Hs.25447	
	GAAGGGGCAAT	1,00			ESTs, Weakly similar to ALU4_HUMAN
		,]	,,		ALU SUBFAMILY SB2
4152	ATGTTGGAAAG	1,00	1.26	Hs.253704	
	CAAATAGGCCA	1,00			hypothetical protein DKFZp761G1923
-	GAGATGTATCT	1,00		Hs.252934	
	GATGACAAAAA	1,00		Hs.252807	
	CCCCAGCAGTC	1,00			ribosomal protein S3
	CCACACAATTC	1,00	1.26		Homo sapiens clone 23967 unknown
		.,00	.,_0		mRNA, partial cds
4158	CAACATTTTAA	1,00	1.26	Hs.250158	
	TATAAAAAGTA	1,00		Hs.249759	
	GGGGCAGATCC	1,00		Hs.249669	
	AGCCACTGTCC	1,00			ESTs, Highly similar to cep250
1	1000,1010100	.,00	.,	110.210.01	centrosome associated
4162	AGCTTTCTCAA	1,00	1 26	Hs.245297	
	TTTCTTTGACC	1,00		Hs.244865	
	GGCCGGGACCC	1,00			EST, Weakly similar to S39803
1.101		1,50	1,20		ribosomal protein L4 -
4165	AGAGATACTAG	1,00	1.26	Hs.244473	
	GAAGCTTGGTC	1,00		Hs.243582	
	GCATCGCTGTT	1,00		Hs.243570	
	GTTAAAACCCC	1,00			EST, Weakly similar to ALU8_HUMAN
		',00	.,_0	10.2.00.2	ALU SUBFAMILY SX S
4169	AAGAGATGTGC	1,00	1.26	Hs.242885	
	TCTACTAAAGA	1,00			EST, Moderately similar to
		',55	.,_0		ALU6_HUMAN ALU SUBFAMILY
4171	GCCACGCCCAG	1,00	1.26	Hs 241507	ribosomal protein S6
	TATAAAACAGA	1,00			ESTs
	GACTITICTGG	1,00		Hs.2407	POU domain, class 2, associating factor
'''		",00	.,	10.2107	1
4174	AAAAACTGTCC	1,00	1 26	Hs 239818	phosphoinositide-3-kinase, catalytic, beta
		.,55	.,20		polypepti
4175	ATAGCTTTGAT	1,00	1.26	Hs.23767	guanine nucleotide binding protein (G
		.,55	.,		protein), gamm
4176	GAATACTACTC	1,00	1.26	Hs.237097	<del></del>
	CCCTTCTTTGT	1,00			hydroxyacid oxidase 2 (long chain)
	TTCTATGATCC	1,00		Hs.235860	<del></del>
	GCACACTCATA	1,00			organic anion transporter OATP-E
1113	JOURDAU TONIA	1,00	1,20	J. 10.2001 02	Joiganno annon transporter OATT-L

4180	GGGACCCTCAG	1,00		Hs.235604	<u> </u>
4181	GATCCCAGTTT	1,00	1,26	Hs.234489	lactate dehydrogenase B
4182	TCCAACAGCCT	1,00	1,26	Hs.233789	ESTs
4183	AAGAAAGACTA	1,00	1,26	Hs.233431	ESTs
4184	GAAGACAAAAG	1,00	1,26	Hs.233383	ESTs
4185	TTGCTTCTCAA	1,00	1,26	Hs.233172	EST
4186	TGTATTTCTTT	1,00	1,26	Hs.233045	EST
4187	GAATAGCTCTG	1,00	1,26	Hs.233041	EST
4188	TGAGCATATTT	1,00	1,26	Hs.232010	ESTs
4189	ATCTAACTAGA	1,00	1,26	Hs.230905	EST
4190	GATCAAAAATA	1,00		Hs.230647	
4191	GGTGCCAGAGT	1,00	1,26	Hs.229501	EST, Weakly similar to ALUC_HUMAN
		•	·		IIIII ALU CLASS C W
4192	GGGTGGGTTTT	1,00	1,26	Hs.229064	
4193	CCTTTGAGATC	1,00	1,26	Hs.228944	EST, Weakly similar to RS5_HUMAN
					40S RIBOSOMAL PROTE
4194	ССТТТСТСТСТ	1,00	1,26	Hs.227948	squamous cell carcinoma antigen 1
4195	TGGCAGCTTTG	1,00		Hs.22708	
4196	TCGACTGAGAA	1,00	1,26	Hs.226133	growth arrest-specific 7
4197	AATCAGGGGAC	1,00			IDN3 protein
4198	TCTTAGTAGAG	1,00		Hs.22564	
4199	CTCCCAGAATC	1,00		Hs.225160	
4200	TGGGGATGAAA	1,00		Hs.225118	
4201	TCCAATAAAAT	1,00	1,26	Hs.224808	EST
4202	TTCCTGAATAA	1,00			EST, Moderately similar to nucleolar
		'	,	-	RNA-helicase [H
4203	CTGAGACTAAA	1,00	1,26	Hs.224015	ESTs
4204	AGCTATAATTG	1,00	1,26	Hs.223572	EST, Weakly similar to ALU1_HUMAN
1 1					ALU SUBFAMILY J SE
4205	AAAGCCAAGAC	1,00	1,26	Hs.222728	ESTs
4206	GCTTTTAGGTT	1,00	1,26	Hs.221660	blood vessel epicardial substance
4207	CAGCACTCCTT	1,00			ESTs, Moderately similar to
			·		ALU1_HUMAN ALU SUBFAMILY
4208	CCTCCCTCGGC	1,00	1,26	Hs.21729	splicing factor 3a, subunit 1, 120kD
4209	CCACTGGAGTC	1,00	1,26	Hs.215893	ESTs, Weakly similar to RMS1_HUMAN
					REGULATOR OF MITO
4210	ATCACCAAAGT	1,00	1,26	Hs.214039	ESTs
4211	CCTCCGGTACT	1,00	1,26	Hs.214004	EST
	GATTCATTGCA	1,00		Hs.214	neuro-oncological ventral antigen 1
4213	AGTCAAGATCA	1,00	1,26	Hs.21321	Homo sapiens mRNA; cDNA
					DKFZp564E1363 (from clone DK
4214	TAAAAAAAGGA	1,00	1,26	Hs.212481	ESTs
4215	CATTGATACTA	1,00	1,26	Hs.210347	ESTs
4216	GAGTCCCGGCT	1,00		Hs.210265	
4217	AGGGGAAGGCG	1,00	1,26	Hs.208035	EST
4218	TGTGATTGTAA	1,00		Hs.207915	
4219	GTAACCAAAAA	1,00		Hs.207593	
4220	ACTITICCCAC	1,00		Hs.207345	
	CACCCTGAATA	1,00		Hs.205769	

	_				
4222	GTGGTTGCCCT	1,00	1,26	Hs.205466	ESTs
	CAAAGTAATCT	1,00		Hs.203800	
	CTTCTGTGTGG	1,00		Hs.202202	
	CTGGCAAATTT	1,00		Hs.202072	
	ACATTTAAAAG	1,00		Hs.20142	
	CACAGCCTGCC	1,00			apyrase, lysosomal
	CTCCCCCCCC	1,00		Hs.199913	
	GTAAGACTCTG	1,00			Homo sapiens mRNA; cDNA
	01,010,01010	',00	1,20	1.0.10002	DKFZp564P1816 (from clone DK
4230	ATTTCTTAGCT	1,00	1 26	Hs 198253	major histocompatibility complex, class
'200		1,00	1,20	110.100200	II, DQ alpha
4231	TATTATGGCCA	1,00	1 26	Hs 197766	Human clone 23932 mRNA sequence
	GGTCAAGTCTG	1,00		Hs.197708	
	TGCAAAAATCT	1,00		Hs.197671	<del></del>
	ATCCATTITGA	1,00			ESTs, Weakly similar to unknown
1201	MOOMITTOA	1,00	1,20	113.13702	[D.melanogaster]
4235	AAGCACCTACG	1,00	1 26	Hs.197143	
	TTATGCTCTTG	1,00		Hs.197143	
	GTGCTCAATTT	1,00		Hs.196663	
	GCGTCTCGATG				<del></del>
		1,00		Hs.196058	
4239	GTAAACGCTGT	1,00	1,26	HS.195175	CASP8 and FADD-like apoptosis
4240	CATTOTOATTO	4 00	4.00	11- 404004	regulator
	CATTCTCATTG	1,00		Hs.194984	<u> </u>
	ACTTGCTGTGT	1,00		Hs.194031	
	TCTGTGTCTAA	1,00		Hs.194024	
	TAACCTAAAAC	1,00			attractin (with dipeptidylpeptidase IV activity)
4244	TTGGTGTCACT	1,00	1,26		ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
4245	CCTCCATCTTC	1,00	1,26		ESTs, Weakly similar to Rigui
			Ţ.		[H.sapiens]
4246	AAAAAGAGTTT	1,00	1,26	Hs.193743	ESTs
4247	TTTAGTGAAAA	1,00	1,26	Hs.193370	ESTs, Highly similar to LIM domains
					containing prote
	TGATCTTTATG	1,00		Hs.192374	
	CCCTCCCAGCT	1,00		Hs.191742	
4250	TTCCCAGCTGC	1,00	1,26	Hs.19121	adaptor-related protein complex 2, alpha 2 subunit
4251	CAAAAACAGTA	1,00	1.26	Hs.191077	
	AGAGCAGGGCA	1,00		Hs.190874	
	GTATGGTAGAG	1,00		Hs.189773	
	TCACCCCTATG	1,00			putative protein similar to nessy
					(Drosophila)
	GAAGCTCCAAA	1,00			hypothetical protein FLJ20070
4256	TATTTTACCGT	1,00	1,26	Hs.184693	transcription elongation factor B (SIII), polypeptid
4257	CGGTCGGGCAG	1,00	1.26	Hs.184634	hypothetical protein FLJ20005
	AGGACATCTAT	1,00			chromosome 2 open reading frame 3
	TCCATAAAGAA	1,00			ankyrin 1, erythrocytic

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A261   GCACTITCCTT   1,00	(45.5		4 2 2			
4262   TGGCTCCCTT			1,00			
type, f polyp   type, f poly						
A263   GGCCAGGTGG	4262	TGGCTCCCCTT	1,00	1,26	Hs.183648	
4264   TCCTACCCTAT						type, f polyp
GUANINĒ NUCLEOTID	4263	GGGCCAGGTGG	1,00	1,26	Hs.183138	procollagen (type III) N-endopeptidase
4265   AGGTTTGAGG	4264	TCCTACCCTAT	1,00	1,26	Hs.181781	
4266   AGTGGAGAGTA						GUANINE NUCLEOTID
MRNA sequence   MRNA sequenc	4265	AGGTTTTGAGG	1,00			
4267   TAAAAGCCTTT	4266	AGTGGAGAGTA	1,00	1,26	Hs.180737	Homo sapiens clone 23664 and 23905
4268   CCCAGGGGAAA						mRNA sequence
4269   GCAAGTTGCCA   1,00   1,26   Hs.180398   LIM domain-containing preferred translocation partne   4270   GTTTTGGTATT   1,00   1,26   Hs.179526   upregulated by 1,25-dihydroxyvitamin E 3   3   4271   GAGATTCTCAT   1,00   1,26   Hs.17917   lymphatic vessel endothelial hyaluronar receptor 1   4272   AATTAGGATAA   1,00   1,26   Hs.17914   ESTs, Weakly similar to AF201951_1   high affinity imm   4273   TCTCCAGGGCC   1,00   1,26   Hs.178803   ESTs   4274   AGCTGTCTTT   1,00   1,26   Hs.177744   ESTs   4275   AGATTTGGGGCG   1,00   1,26   Hs.1777455   ESTs   4276   TCCCTTTTC   1,00   1,26   Hs.177765   ESTs   4276   TCCCTTTTTC   1,00   1,26   Hs.177765   ESTs   4279   TATGTCTTGGA   1,00   1,26   Hs.1777162   ESTs   4279   TATGTCTTGGA   1,00   1,26   Hs.177096   ESTs   4279   TATGTCTTGGA   1,00   1,26   Hs.17605   Homo sapiens mRNA full length insert   kinase   4280   CAGTAAACCTC   1,00   1,26   Hs.175563   Homo sapiens mRNA; cDNA   DKFZp564N0763 (from clone DK   4281   TGGATGAAAAA   1,00   1,26   Hs.175551   EST   4284   GAAGTTCTGC   1,00   1,26   Hs.175339   EST   4284   GAAGTTCTGC   1,00   1,26   Hs.175339   EST   4285   GGGAAAAAAAA   1,00   1,26   Hs.175350   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175351   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175359   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175359   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175359   EST   4286   ACCAAAGAGAC   1,00   1,26   Hs.175359   transforming, acidic colled-coil containing protein   4288   ACCAAAGAGAC   1,00   1,26   Hs.172652   KlAA0013 gene product   4289   AAAGGAAGATT   1,00   1,26   Hs.172652   KlAA0013 gene product   4289   AAAGGAAGATT   1,00   1,26   Hs.172675   right functional domain (PTPRF interacting)   4292   GATATGGTTTG   1,00   1,26   Hs.170773   ESTs   4292   GATATGGTTTG   1,00   1,26   Hs.17077	4267	TAAAAGCCTTT	1,00	1,26	Hs.180668	ESTs
4269   GCAAGTTGCCA   1,00   1,26   Hs.180398   LIM domain-containing preferred translocation partne   4270   GTTTTGGTATT   1,00   1,26   Hs.179526   upregulated by 1,25-dihydroxyvitamin E 3   3   4271   GAGATTCTCAT   1,00   1,26   Hs.17917   lymphatic vessel endothelial hyaluronar receptor 1   4272   AATTAGGATAA   1,00   1,26   Hs.17914   ESTs, Weakly similar to AF201951_1   high affinity imm   4273   TCTCCAGGGCC   1,00   1,26   Hs.178803   ESTs   4274   AGCTGTCTTT   1,00   1,26   Hs.177744   ESTs   4275   AGATTTGGGGCG   1,00   1,26   Hs.1777455   ESTs   4276   TCCCTTTTC   1,00   1,26   Hs.177765   ESTs   4276   TCCCTTTTTC   1,00   1,26   Hs.177765   ESTs   4279   TATGTCTTGGA   1,00   1,26   Hs.1777162   ESTs   4279   TATGTCTTGGA   1,00   1,26   Hs.177096   ESTs   4279   TATGTCTTGGA   1,00   1,26   Hs.17605   Homo sapiens mRNA full length insert   kinase   4280   CAGTAAACCTC   1,00   1,26   Hs.175563   Homo sapiens mRNA; cDNA   DKFZp564N0763 (from clone DK   4281   TGGATGAAAAA   1,00   1,26   Hs.175551   EST   4284   GAAGTTCTGC   1,00   1,26   Hs.175339   EST   4284   GAAGTTCTGC   1,00   1,26   Hs.175339   EST   4285   GGGAAAAAAAA   1,00   1,26   Hs.175350   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175351   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175359   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175359   EST   4286   CTACCTGACCT   1,00   1,26   Hs.175359   EST   4286   ACCAAAGAGAC   1,00   1,26   Hs.175359   transforming, acidic colled-coil containing protein   4288   ACCAAAGAGAC   1,00   1,26   Hs.172652   KlAA0013 gene product   4289   AAAGGAAGATT   1,00   1,26   Hs.172652   KlAA0013 gene product   4289   AAAGGAAGATT   1,00   1,26   Hs.172675   right functional domain (PTPRF interacting)   4292   GATATGGTTTG   1,00   1,26   Hs.170773   ESTs   4292   GATATGGTTTG   1,00   1,26   Hs.17077	4268	CCCAGGGGAAA		1,26	Hs.180414	heat shock 70kd protein 10 (HSC71)
translocation partne   translocation   translocation partne   translocation   translocati						
4270   GTTTTGGTATT			.,	.,		
3   3   3   3   3   3   3   3   3   3	4270	GTTTTGGTATT	1.00	1.26		
A272   AATTAGGATAA			.,00	.,_0		3
A272   AATTAGGATAA	4271	GAGATTCTCAT	1.00	1.26	Hs.17917	lymphatic vessel endothelial hyaluronan
4272   AATTAGGATAA			",""	.,_0		
high affinity imm	4272	AATTAGGATAA	1.00	1.26		
4273 TCTCCAGGGCC         1,00         1,26 Hs.178803 ESTs           4274 AGCTGTCTCTT         1,00         1,26 Hs.177744 ESTs           4276 TCCCTTTTC         1,00         1,26 Hs.177655 ESTs           4276 TCCCTTTTC         1,00         1,26 Hs.177486 amyloid beta (A4) precursor protein (protease nexin-qrotease			.,	,,_0		
4274 AGCTGTCTCTT         1,00         1,26 Hs.177744 ESTs           4275 AGATTTGGGCG         1,00         1,26 Hs.177655 ESTs           4276 TTCCCTTTTC         1,00         1,26 Hs.177655 ESTs           4277 GGAGTGCAAAT         1,00         1,26 Hs.177162 ESTs           4278 GTTGTTTTTA         1,00         1,26 Hs.17796 ESTs           4279 TATGTCTTGGA         1,00         1,26 Hs.17605 Iymphocyte-specific protein tyrosine kinase           4280 CAGTAAACCTC         1,00         1,26 Hs.176005 Homo sapiens mRNA full length insert cDNA clone EURO           4281 TGGATGAAAAA         1,00         1,26 Hs.175563 Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAA         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175350 EST           4284 GAAGTTTCTGC         1,00         1,26 Hs.175339 EST           4285 GGGAAGTTCTT         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4286 CTACCTGACCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4287 GAAAGGACCCT         1,00         1,26 Hs.172652 KIAA0013 gene product           4288 ACCAAAGAGAC         1,00         1,26 Hs.172185 cell division cycle 27           4290 CTTATATATCTC         1,00         1,26 Hs.172195 cell division domain (PTPRF	4273	TCTCCAGGGCC	1.00	1.26	Hs 178803	
4275 AGATTTGGGCG         1,00         1,26 Hs.177655 ESTs           4276 TTCCCTTTTC         1,00         1,26 Hs.177486 amyloid beta (A4) precursor protein (protease nexin-4277 GGAGTGCAAAT         1,00         1,26 Hs.177162 ESTs           4278 GTTGTTTTTA         1,00         1,26 Hs.177096 ESTs         lymphocyte-specific protein tyrosine kinase           4280 CAGTAAACCTC         1,00         1,26 Hs.17650 Hs.176005 Homo sapiens mRNA full length insert cDNA clone EURO           4281 TGGATGAAAAA         1,00         1,26 Hs.175563 Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAA         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175339 EST           4284 GAAGTTCTG         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4285 GGGAAGTTCTT         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4286 CTACCTGACCT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGACCT         1,00         1,26 Hs.172652 KIAA0013 gene product           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interac						
4276         TTCCCTTTTC         1,00         1,26         Hs.177486 amyloid beta (A4) precursor protein (protease nexin-4277 GGAGTGCAAAT         1,00         1,26         Hs.177162 ESTs           4278         GTTGTTTTTA         1,00         1,26         Hs.177096 ESTs         lymphocyte-specific protein tyrosine kinase           4280         CAGTAAACCTC         1,00         1,26         Hs.176005 Homo sapiens mRNA full length insert cDNA clone EURO           4281         TGGATGAAAAA         1,00         1,26         Hs.175563 Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282         GGGAAAAAAAG         1,00         1,26         Hs.175351 EST           4283         ATTGTGGGGCC         1,00         1,26         Hs.175350 EST           4284         GAAGTTCTGC         1,00         1,26         Hs.175339 EST           4285         GGGAAGTTCTT         1,00         1,26         Hs.175218 potassium channel, subfamily K, memb 7           4286         CTACCTGACCT         1,00         1,26         Hs.173159 transforming, acidic coiled-coil containing protein           4287         GAAAGGGCCCT         1,00         1,26         Hs.172652 KIAA0013 gene product           4289         AAAGGAAGATT         1,00         1,26         Hs.172182 poly(A)-binding protein, cytoplasmic 1						
(protease nexin-    4277 GGAGTGCAAAT   1,00   1,26   Hs.177162   ESTs     4278 GTTGTTTTTA   1,00   1,26   Hs.177096   ESTs     4279 TATGTCTTGGA   1,00   1,26   Hs.1765     1,00   1,26   Hs.176005     1,00   1,26   Hs.176005     1,00   1,26   Hs.175005     1,00   1,26   Hs.175005     1,00   1,26   Hs.175503     1,00   1,26   Hs.175503     1,00   1,26   Hs.175503     1,00   1,26   Hs.175351   EST   1,00   1,26   Hs.175350   EST   1,00   1,26   Hs.175350   EST   1,00   1,26   Hs.175339   EST   1,00   1,26   Hs.175218   1,00   1,26   Hs.175218   1,00   1,26   Hs.175218   1,00   1,26   Hs.173159   1,00   1,26   Hs.172182   1,00   1,26   Hs.172182   1,00   1,26   Hs.172405   1,00   1,26						
4277 GGAGTGCAAAT         1,00         1,26 Hs.177162 ESTs           4278 GTTGTTTTTA         1,00         1,26 Hs.177096 ESTs           4279 TATGTCTTGGA         1,00         1,26 Hs.1765   lymphocyte-specific protein tyrosine kinase           4280 CAGTAAACCTC         1,00         1,26 Hs.176005   Homo sapiens mRNA full length insert cDNA clone EURO           4281 TGGATGAAAAA         1,00         1,26 Hs.175563   Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAG         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175339 EST           4284 GAAGTTCTGC         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4285 GGGAAGGCCT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs			.,00	1,20		
4278 GTTGTTTTTA         1,00         1,26 Hs.177096 ESTs           4279 TATGTCTTGGA         1,00         1,26 Hs.1765         lymphocyte-specific protein tyrosine kinase           4280 CAGTAAACCTC         1,00         1,26 Hs.176005         Homo sapiens mRNA full length insert cDNA clone EURO           4281 TGGATGAAAAA         1,00         1,26 Hs.175563         Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAG         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175339 EST           4284 GAAGTTCTT         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4286 CTACCTGACCT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 irriple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs	4277	GGAGTGCAAAT	1.00	1 26		
4279 TATGTCTTGGA         1,00         1,26 Hs.1765         lymphocyte-specific protein tyrosine kinase           4280 CAGTAAACCTC         1,00         1,26 Hs.176005         Homo sapiens mRNA full length insert cDNA clone EURO           4281 TGGATGAAAAA         1,00         1,26 Hs.175563         Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAG         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175350 EST           4284 GAAGTTCTGC         1,00         1,26 Hs.175318 potassium channel, subfamily K, memb 7           4285 GGGAAGTCTT         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4286 CTACCTGACCT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs						
Kinase   K						
4280 CAGTAAACCTC         1,00         1,26 Hs.176005         Homo sapiens mRNA full length insert cDNA clone EURO           4281 TGGATGAAAAA         1,00         1,26 Hs.175563         Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAG         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175339 EST           4284 GAAGTTCTG         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4285 GGGAAGTCTT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.171957 interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs	12,0	////C/01/00/	1,00	1,20		
CDNA clone EURO	4280	CAGTAAACCTC	1.00	1 26		
4281 TGGATGAAAAA         1,00         1,26 Hs.175563 Homo sapiens mRNA; cDNA DKFZp564N0763 (from clone DK           4282 GGGAAAAAAAG         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175350 EST           4284 GAAGTTCTGC         1,00         1,26 Hs.175339 EST           4285 GGGAAGTTCTT         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4286 CTACCTGACCT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs		0/101/1110010	',"	1,20		
DKFZp564N0763 (from clone DK	4281	TGGATGAAAA	1 00	1 26		
4282 GGGAAAAAAAG         1,00         1,26 Hs.175351 EST           4283 ATTGTGGGGCC         1,00         1,26 Hs.175350 EST           4284 GAAGTTCTGC         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4285 GGGAAGTTCTT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs			1,00	1,20	110.170000	
4283 ATTGTGGGGCC         1,00         1,26 Hs.175350 EST           4284 GAAGTTCTGC         1,00         1,26 Hs.175339 EST           4285 GGGAAGTTCTT         1,00         1,26 Hs.175218 potassium channel, subfamily K, memb 7           4286 CTACCTGACCT         1,00         1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)           4287 GAAAGGGCCCT         1,00         1,26 Hs.173159 transforming, acidic coiled-coil containing protein           4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs	4282	GGGAAAAAAG	1.00	1 26	He 175351	
4284 GAAGTTTCTGC1,001,26 Hs.175339 EST4285 GGGAAGTTCTT1,001,26 Hs.175218 potassium channel, subfamily K, memb 74286 CTACCTGACCT1,001,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)4287 GAAAGGGCCCT1,001,26 Hs.173159 transforming, acidic coiled-coil containing protein4288 ACCAAAGAGAC1,001,26 Hs.172652 KIAA0013 gene product4289 AAAGGAAGATT1,001,26 Hs.172405 cell division cycle 274290 CTTATAATCTC1,001,26 Hs.172182 poly(A)-binding protein, cytoplasmic 14291 AAGTATCAGCT1,001,26 Hs.171957 triple functional domain (PTPRF interacting)4292 GATATGGTTTG1,001,26 Hs.170773 ESTs						
4285 GGGAAGTTCTT 1,00 1,26 Hs.175218 potassium channel, subfamily K, memb 7  4286 CTACCTGACCT 1,00 1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)  4287 GAAAGGGCCCT 1,00 1,26 Hs.173159 transforming, acidic coiled-coil containing protein  4288 ACCAAAGAGAC 1,00 1,26 Hs.172652 KIAA0013 gene product 4289 AAAGGAAGATT 1,00 1,26 Hs.172405 cell division cycle 27 4290 CTTATAATCTC 1,00 1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1 4291 AAGTATCAGCT 1,00 1,26 Hs.171957 triple functional domain (PTPRF interacting)  4292 GATATGGTTTG 1,00 1,26 Hs.170773 ESTs						
4286 CTACCTGACCT 1,00 1,26 Hs.174044 dishevelled 3 (homologous to Drosophil dsh)  4287 GAAAGGGCCCT 1,00 1,26 Hs.173159 transforming, acidic coiled-coil containing protein  4288 ACCAAAGAGAC 1,00 1,26 Hs.172652 KIAA0013 gene product  4289 AAAGGAAGATT 1,00 1,26 Hs.172405 cell division cycle 27  4290 CTTATAATCTC 1,00 1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1  4291 AAGTATCAGCT 1,00 1,26 Hs.171957 triple functional domain (PTPRF interacting)  4292 GATATGGTTTG 1,00 1,26 Hs.170773 ESTs						
dsh	7203	GGGAAGIICII	1,00	1,20	ПS. 1752 16	potassium channel, subtamily K, member
dsh	4286	CTACCTGACCT	1.00	1 26	Uc 174044	dishavalled 2 (hamalagava ta Draganhila
4287GAAAGGGCCCT1,001,26Hs.173159transforming, acidic coiled-coil containing protein4288ACCAAAGAGAC1,001,26Hs.172652KIAA0013 gene product4289AAAGGAAGATT1,001,26Hs.172405cell division cycle 274290CTTATAATCTC1,001,26Hs.172182poly(A)-binding protein, cytoplasmic 14291AAGTATCAGCT1,001,26Hs.171957triple functional domain (PTPRF interacting)4292GATATGGTTTG1,001,26Hs.170773ESTs	7200	O I AGG I GAGG I	1,00	1,20	П <b>5.</b> 1740 <del>44</del>	
Protein	4297	GAAAGGCCCT	1.00	1.26	Un 172150	
4288 ACCAAAGAGAC         1,00         1,26 Hs.172652 KIAA0013 gene product           4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs	4201	GAAAGGGCCC1	1,00	1,20	ns. 173159	
4289 AAAGGAAGATT         1,00         1,26 Hs.172405 cell division cycle 27           4290 CTTATAATCTC         1,00         1,26 Hs.172182 poly(A)-binding protein, cytoplasmic 1           4291 AAGTATCAGCT         1,00         1,26 Hs.171957 triple functional domain (PTPRF interacting)           4292 GATATGGTTTG         1,00         1,26 Hs.170773 ESTs	1200	ACCAAACACAC	1.00	4.26	Un 470650	
4290 CTTATAATCTC1,001,26 Hs.172182 poly(A)-binding protein, cytoplasmic 14291 AAGTATCAGCT1,001,26 Hs.171957 triple functional domain (PTPRF interacting)4292 GATATGGTTTG1,001,26 Hs.170773 ESTs						
4291 AAGTATCAGCT 1,00 1,26 Hs.171957 triple functional domain (PTPRF interacting) 4292 GATATGGTTTG 1,00 1,26 Hs.170773 ESTs						
interacting)   4292 GATATGGTTTG   1,00   1,26 Hs.170773 ESTs				1,26	ms.1/2182	poly(A)-binding protein, cytoplasmic 1
4292 GATATGGTTTG 1,00 1,26 Hs.170773 ESTs	4291	ANGIATUAGUT	1,00	1,26		
	4200	CATATOOTTO	-4-00	4.00		
4293 TAACTCTAGAA   1,00   1,26 Hs.170714 ESTs	4233	IMACICIAGAA	1,00	1,20	ms.1/0/14	EOIS

PCT/EP01/15179

1.55					
4294	AGAAGATGCCG	1,00	1,26	Hs.169902	solute carrier family 2 (facilitated glucose
4005	TTT00040444	4.00	4.00		transpo
	TTTCGCAGAAA	1,00		Hs.169395	
	CTTTTGTAATG	1,00			karyopherin alpha 1 (importin alpha 5)
	AAATGATCCCG	1,00		Hs.168941	
	TTCAAAGTTGA	1,00		Hs.168363	
	TAATTGTTTAT	1,00			hypothetical protein FLJ10242
	GGCAAGTTCCT	1,00		Hs.166832	
4301	GGAGATAGTGC	1,00	1,26	Hs.166369	ESTs, Highly similar to sodium
					bicarbonate cotranspo
	TTCTGCTTTCG	1,00		Hs.1652	chemokine (C-C motif) receptor 7
	TCACGCCGGAC	1,00		Hs.165003	
	TTCCCTTTATT	1,00		Hs.164427	
4305	AATATGTACAC	1,00	1,26	Hs.163900	ESTs
4306	CAAGAACAATC	1,00	1,26	Hs.163724	HSPC019 protein
4307	GGCAGCTGGTA	1,00		Hs.163252	
4308	AGGGATATTGG	1,00	1,26	Hs.163203	ESTs, Weakly similar to B34087
			•		hypothetical protein
4309	ACACTGTACTC	1,00	1.26	Hs.163189	
	GAAGCAGAAAA	1,00		Hs.162852	
	GAGCAAACTTT	1,00		Hs.16249	
	GACTGGAAGTA	1,00		Hs.162252	
	AGGCTGAGGAA	1,00		Hs.161790	
	TGGAAACTCAG	1,00		Hs.161719	
	ACAGATTTTGG	1,00			hypothetical protein FLJ20159
	ATTCTTTAAAA	1,00			hypothetical protein FLJ20159
	CTAGTGCACAC	1,00			hypothetical protein FLJ20159
	GGCAACAAGGT	1,00			hypothetical protein FLJ20159
	TAAACCTTTTG	1,00			hypothetical protein FLJ20159
	TTGAAATTTGA	1,00			tachykinin receptor 2
	GTCTGTGCTAT	1,00		Hs.161277	
	TTTCTTGGTAA	1,00		Hs.161065	
	AAAAATGCTTC	1,00		Hs.160914	
	CTAAATTCGGA			Hs. 160275	
		1,00			
	CCAGCTAGAAA	1,00		Hs.160013	
	GAGCAGCTCAC	1,00			chemokine-like receptor 1
	TAGAGTGATGG	1,00		Hs.159153	
	TACCCAGACTT	1,00			KIAA0468 gene product
4329	CCACCTGAATT	1,00	1,26	Hs.15740	Homo sapiens mRNA; cDNA
1000					DKFZp434E033 (from clone DKF
	GGGCTGGTGGA	1,00		Hs.156836	
	TCACGAGCTAT	1,00		Hs.155764	
4332	CCCTTTGAGCA	1,00	1,26	Hs.155381	ESTs, Moderately similar to I38022
10.00					hypothetical prot
	GAATTTACTTA	1,00			KIAA0095 gene product
4334	TCAGGGTGAAG	1,00	1,26	Hs.154156	myosin, light polypeptide 4, alkali; atrial,
					embryon
	GGACTTTCAAA	1,00		Hs.153739	
4336	AGAGAAGATGC	1,00	1,26	Hs.153716	Homo sapiens mRNA for Hmob33

		Т			protein, 3' untranslate
1227	TACTCCCCTGA	1,00	1 26	Un 152026	adaptor-related protein complex 2, mu 1
4331	IACICCCIGA	1,00	1,20	ns. 152930	Isubunit
1220	TOACCTACTOC	1,00	1 26	Hs.152666	
	TGACCTACTGG AGTGTAAGTTT				
		1,00		Hs.152129	
	CACCGTGTGTC	1,00		Hs.152016	
	TAACTCCTCCC	1,00		Hs.150784	
4342	GGGTAAGGAGG	1,00	1,26	HS. 150423	cyclin-dependent kinase 9 (CDC2-related
4242	AATTTATATAA	1.00	4.26	Un 450072	kinase) ESTs, Moderately similar to alternatively
4343	MITIATATAA	1,00	1,20		
1311	TGTAATGTAAA	1.00	1 26	He 140622	spliced pr group III secreted phospholipase A2
	TGGGTGAAAAA	1,00		Hs.148725	
	TTGTCTACAAA	1,00			ESTs, Moderately similar to AF161544_1
4340	ITGICIACAAA	1,00	1,20	ns. 140217	
1317	AACTTGGATAT	1,00	1 26	Hs.148129	HSPC059 [H.sa
	TTAACTTTTTT	1,00		Hs.148091	
	TTTCTTAGGTT	1,00		Hs.147317	
	CCTCCCCCATT			Hs.146893	
	TTGTTGCTGGT	1,00			
		1,00		Hs.146730	
	GTGCTGTTAGT	1,00		Hs.146712	
	ATGACATTGGA	1,00		Hs.146335	
4354	ATTGGTAGACA	1,00	1,20	HS. 146311	ESTs, Moderately similar to
1255	TITATOTITI	4.00	1.00	11- 445504	ALU1_HUMAN ALU SUBFAMILY
	TTTATCTTTT	1,00		Hs.145584	
4330	TTGTCAAAAAA	1,00	1,26	HS. 1452/9	SET translocation (myeloid leukemia-
1257	TATATTATAGA	1,00	1.26	Un 144021	associated) ATPase, aminophospholipid transporter
4337	IAIAIIAIAGA	1,00	1,20	П5. 14493 І	(APLT), Class
1358	GGAATTGTTGT	1,00	1 26	Hs.143866	
	AGCCTTTGCCA	1,00		Hs.143574	
	TCTTTATTTTC	1,00			flavin containing monooxygenase 5
	TTTCCTATAAA			Hs.141939	
	TCTAAAGAATT	1,00			Homo sapiens clone 23758 mRNA
4302	TOTAMOMIT	1,00	1,20	HS. 14 1000	sequence
4363	TCAGCGGACTG	1,00	1 26	He 138280	ESTs, Moderately similar to
7505	10000000010	1,00	1,20	118.130200	ALU1_HUMAN ALU SUBFAMILY
4364	ACAATGTAGGA	1,00	1 26	He 137415	Homo sapiens BAC clone RP11-294L11
1001	10,000	1,00	1,20	113.137-13	from 2
4365	TGACTTGGCCA	1,00	1 26	Hs.137401	
	CTCCCCAAAAA	1,00			hypothetical protein FLJ10761
$\overline{}$	CTCCCTCTTTC	1,00		Hs.136468	
	GAAGCCCAATT	1,00		Hs.136432	<del></del>
	CTCTAGGTAAA	1,00	_	Hs.136423	<u> </u>
	GTGATATGACT	1,00		Hs.136200	
	GAGAGAAATGG	1,00			Homo sapiens clone 25215 mRNA
' '	-, 10, 10, 17, 100	1,00	1,20	1 13. 100 103	sequence, partial cds
4372	CACATTTTAA	1,00	1 26	Hs.136008	
	GTTTTCTCTGT	1,00		Hs.135503	
		.,00	1,20		

4374	TTCATAAGTTT	1,00	1,26	Hs.135473	ESTs
4375	AGACGGTCCAA	1,00	_1,26	Hs.134757	hypothetical protein FLJ20033
4376	TACCCCAGGAA	1,00	1,26	Hs.134350	ESTs
4377	TTGCTCCTCCT	1,00	1,26	Hs.133521	ESTs
4378	CAGGGGAGGAC	1,00		Hs.133386	
4379	AATATGCGGCA	1,00	1,26	Hs.133355	ESTs
4380	AGTTTGAGATA	1,00	1,26	Hs.133266	ESTs
	AATCTGGCGTA	1,00			ESTs, Moderately similar to p53
1		, ,			regulated PA26-T2 nu
4382	GTCAGGAGAAG	1,00	1,26	Hs.132278	
	TCTTTATTTTT	1,00		Hs.131360	
	GCCTGCAGGAA	1,00		Hs.131030	
	TTGATTCCTAT	1,00			hypothetical protein FLJ10173
	TTTAGAGTGGA	1,00		Hs.130716	
	TGTCTGGTTTA	1,00		Hs.130518	
	TTCATTCATTC	1,00		Hs.130253	
	GGAGGGCTGTG	1,00			ESTs, Moderately similar to melastatin 1
		1,55	.,		[H.sapiens]
4390	CTTCCTGCTAC	1,00	1.26	Hs.128018	
	TCTAAGCTTGT	1,00		Hs.127780	
	TGAGAATTCTG	1,00		Hs.127378	
	GTGCCTCAGCC	1,00		Hs.126943	
	CCCCAAGGCCT	1,00			Thy-1 cell surface antigen
	AGGGAGGCTGA	1,00		Hs.125246	
$\overline{}$	TITIGTCTGTCT	1,00			KIAA1172 protein
	CACAGCCACTA	1,00		Hs.125103	
	GATGCTAGAGC	1,00		Hs.124597	
	ATCTTCAGTAT	1,00			CD9 antigen (p24)
	GTCTCCAAGCC	1,00		Hs.124367	
	CATTTCCCTTG	1,00		Hs.123307	
	ATAAAAGGAAA	1,00		Hs.122236	
	ATGAAGTGTTG	1,00			fer (fps/fes related) tyrosine kinase
	7.10,7.010110	1,00	1,20	113.121000	(phosphoprotei
4404	TGTTACTGGGA	1,00	1 26	Hs.120592	
	CCACAGGGATT	1,00			collagen, type III, alpha 1 (Ehlers-Danlos
		.,00	1,20	1.0.770077	syndrome
4406	TTCCTGGTGCG	1,00	1 26	Hs 119251	ubiquinol-cytochrome c reductase core
}		.,00	,,_0	110.110201	protein I
4407	CTAGTTTAAAA	1,00	1 26	Hs.11899	3-hydroxy-3-methylglutaryl-Coenzyme A
		',55	1,20	1 10.1 1000	reductase
4408	ATTAAGTACAA	1,00	1.26	Hs.118959	
	CACTTGTTTCT	1,00			annexin A6
	CCCCGTGGTCA	1,00			ESTs, Highly similar to unnamed protein
		.,55	1,20		product [H.s
4411	CTATAATCCCA	1,00	1 26	Hs 117582	CGI-43 protein
	GCTAAACTCCG	1,00			CGI-43 protein
	GGGTCTTGGAG	1,00			CGI-43 protein
	GTGGCTTTCGC	1,00			CGI-43 protein
	TTTATTGTAGA	1,00			CGI-43 protein
1.7.0	- TINITOTAGA	1,00	1,20	<u>   13.117.002</u>	DOI-TO PROBIN

4416	TGGACAGTGAA	1,00	1,26	Hs.116823	EST
4417	TAACCTAGATG	1,00		Hs.116486	
4418	ACTGCCTGCAT	1,00		Hs.116192	
	GATAAACCAGA	1,00		Hs.115580	
	TTTTACAAACC	1,00	1.26	Hs.115175	sterile-alpha motif and leucine zipper
		","	-,		containing ki
4421	TGTATTGTAAA	1,00	1.26	Hs.115129	
	AAGGGATGCTG	1,00			small inducible cytokine subfamily A
		1,	.,		(Cys-Cys), memb
4423	GGTAAAAAATG	1,00	1.26	Hs.112264	
-	AAAGCCCAAGT	1,00		Hs.111758	
	CAGGGTGTGTG	1,00			ESTs, Moderately similar to dJ108K11.3
		.,,,,,	,		[H.sapiens]
4426	TTTGGCTTGGC	1,00	1 26	Hs.110028	
	AAACAGTAAAT	1,00			zinc finger protein 198
	CCTCATTTCCC	1,00			DKFZP586P2421 protein
	TTGTTGATCCA	1,00			ATP synthase, H+ transporting,
		1,00	.,20	110.107170	mitochondrial F1F0, s
4430	TTTGAAAACAA	1,00	1.26	Hs 107203	hypothetical protein from EUROIMAGE
		.,00	.,20		1759349
4431	AACCTTCCATA	1,00	1.26	Hs.106833	
	AGGAGCACCGT	1,00		Hs.106432	
	AGCTCCGGGAC	1,00			protein phosphatase 1, regulatory
00		1,55	.,_0	110.100010	subunit 10
4434	ATGTTTACCAG	1,00	1 26	Hs.105965	
	ATACGTTTTTT	1,00			KIAA0620 protein
-	ATGACATCACT	1,00		Hs.105506	
-	CCCCTGCCCTG	1,00		Hs.104866	
	ATGTAGTCATT	1,00		Hs.104219	
	TTTCTGCCTTC	1,00		Hs.1042	Sjogren syndrome antigen A1 (52kD,
1700	11101000110	1,00	1,20	115.1042	ribonucleoprotein
4440	CAAATGTCCAT	1,00	1 26	Hs.10359	ESTs
	GCGGTGTACAC	1,00			natural killer cell group 7 sequence
	TACAAAAAAA	1,00			Human DNA sequence from clone RP11-
7742		1,00	0, 14	118.90500	39402 on chromoso
4443	GGACAAAAAAG	1,00	0.14	Hs.84746	
	CGAAGGCTGTA	1,00	0, 14	He 70334	chromosome condensation 1 nuclear factor, interleukin 3 regulated
	ATTGTAAGTTT	1,00	0,14	Hs.74649	cytochrome c oxidase subunit VIc
	GCCACACTGTC	1,00		Hs.61763	ESTs
	CTTCAAGGCCG	_		Hs.59457	ESTs ESTS
<del></del>	CCACACCTCTC	1,00		Hs.54673	
7740	CONCACCICIC	1,00	0,14	⊓S.340/3	tumor necrosis factor (ligand)
4440	GACATCTCTTC	1.00	0.44	Uc 42640	superfamily, member 1 ESTs
		1,00			
	CCTGGCCCTTA	1,00		Hs.42287	E2F transcription factor 6
4401	AAGGGCCACAA	1,00	U, 14	Hs.37096	zinc finger protein 145 (Kruppel-like,
AAEC	TITATITAAT	4 60	044	U- 07040	expressed in
4452	TTATTTTAAT	1,00	0,14	Hs.37040	platelet-derived growth factor alpha
1152	CTITCTCAAAT	4.00	0.44	Un 20507	polypeptide
4433	GTTTCTCAAAT	1,00	0,14	Hs.32597	ring finger protein (C3H2C3 type) 6

T 4 2 4		4 6 6 1	2.4.4		100000000000000000000000000000000000000
	AAATTGCTTAG	1,00			KIAA1344 protein
	GAAAAAGATGT	1,00			AD-017 protein
4456	TTTCAAATAAA	1,00	0,14	Hs.272023	transforming, acidic coiled-coil containing protein
4457	TGGGGAATAGG	1,00	0,14	Hs.23964	sin3-associated polypeptide, 18kD
4458	GTGCAAGCTGT	1,00	0,14	Hs.232068	transcription factor 8 (represses
					interleukin 2 expr
4459	CCTGGGGTAAG	1,00	0,14	Hs.198253	major histocompatibility complex, class
					II, DQ alpha
4460	TGTGAGCCTCA	1,00			cyclin F
4461	GAGTTGGCACT	1,00	0,14	Hs.194688	bromodomain adjacent to zinc finger domain, 1B
4462	GACTGGAACTT	1,00	0,14	Hs.186756	KIAA1441 protein
4463	GCGGCAAGCAT	1,00	0,14	Hs.182729	hypothetical protein FLJ10581
4464	TGGATCACCAA	1,00	0,14	Hs.17872	ESTs
4465	GGAGGGGTTCA	1,00	0,14	Hs.173088	ESTs
4466	CCTTATGGAAA	1,00		Hs.17118	ESTs, Weakly similar to B0025.2 [C.elegans]
4467	GTGGCAGTGGC	1,00	0.14	Hs.16478	
	TGTCCGTCACA	1,00			E74-like factor 1 (ets domain
1 1		, , ,	•		transcription factor)
4469	AAATGGCCAAC	1,00	0,14		DKFZP434J214 protein
$\overline{}$	TACTGTTTGAT	1,00		Hs.12313	
	CGTAAGACGTT	1,00			B-cell CLL/lymphoma 9
4472	GGCTTTCAGCA	1,00			CD59 antigen p18-20 (antigen identified
			,		by monoclona
4473	TAACCCCAAAT	1,00	0,14	Hs.11493	
4474	GTCAAAATTTC	1,00			thrombospondin 2
4475	стттстстс	1,00	0,14	Hs.106823	H.sapiens gene from PAC 426I6, similar to syntaxin 7
4476	CTATTCTAAAA	1,00	0,14		KIAA0692 protein
4477	GTGTCCTCCTC	4,00			Golgi apparatus protein 1
4478	TTGGCTTTTCT	13,00			hypothetical protein
4479	AAGGGGCAAG	6,00			integrin, beta 4
4480	GTGGCTCATAC	3,00	0,35	Hs.281094	ESTs
	GTAACAAGCTC	3,00			KIAA0438 gene product
4482	GACTCAGGGAT	3,00			GTP binding protein 2
	AGGGACATAAA	3,00			BAI1-associated protein 3
	CCTGTGTGTGT	5,00			hypothetical protein FLJ10305
	GGCTGGTCACC	2,00			ESTs, Highly similar to WWP2
					[H.sapiens]
4486	GACATTTTTCC	2,00	0,27	Hs.8083	hypothetical protein FLJ10769
4487	GGGAGGAGGTT	2,00			ESTs
4488	TTTTGTGCATT	2,00	0,27	Hs.238990	cyclin-dependent kinase inhibitor 1B (p27, Kip1)
4489	TGCAATAGGGA	2,00	0 27	Hs 235975	hypothetical protein DKFZp434D0412
	TGGTAGCAGTA	2,00			hypothetical protein FLJ10788
-	GGCTTGGGGAG	2,00			hypothetical protein FLJ10120
	ACTCCAAAAAA	54,00			Homo sapiens mRNA; cDNA
		,00	<u> </u>	1. 10. 1020 10	i oapiono iii tiwi, obiwi

	1			Γ	DKFZp564H172 (from clone DKF
4493	TGGGGAGAGGA	16,00	0.71	Hs.75799	protease, serine, 8 (prostasin)
	TGCTTGACAAG	3,00			RNA polymerase I 16 kDa subunit
	TAGGATGGGG	8,00		Hs.76941	ATPase, Na+/K+ transporting, beta 3
1,100	1710071100000	0,00	0,00	113.700-71	polypeptide
4496	TAGGTTGTCTA	114,00	1 14	Hs 279860	hypothetical protein FLJ20030
_	TAAACTGAAAA	4,00		Hs.3491	ribosomal protein S14
	AAAACATTATG	2,00		Hs.80917	adaptor-related protein complex 3, sigma
	,	-,55	٠,		11 subunit
4499	AATGTCCAGTA	2,00	0.27	Hs.26373	ESTs, Moderately similar to
		'	·		ALU1_HUMAN ALU SUBFAMILY
4500	GTGAAACTGCA	2,00	0,27	Hs.237055	EST
4501	GCACAGATTAC	2,00	0,27	Hs.158497	KIAA0724 gene product
4502	GCACCTGTCGC	2,00		Hs.1239	alanyl (membrane) aminopeptidase
				·	(aminopeptidase N,
	CTGGTGAGTGC	2,00			MYLE protein
4504	ATGATGATGAT	9,00	0,59	Hs.79172	solute carrier family 25 (mitochondrial
4505	0010000000				carrier; ade
	GGAGCGTGGGG	7,00			cathepsin B
	CTCCCCCAAAA	5,00	0,46	Hs.283305	immunoglobulin heavy contant alpha 1
$\overline{}$	AAAACCTGTAA	3,00			coated vesicle membrane protein
4508	TGGCAGTCTGC	3,00	0,36	Hs.6179	DEAD/H (Asp-Glu-Ala-Asp/His) box
4500	TTGTGAGAATA	2.00	0.36	LIA 47740E	polypeptide 17 (72k
	AAGGCGTTTCC	3,00			KIAA0964 protein
	ATACTGTCAGT	3,00	0,36		KIAA0930 protein
	CCTATTTACTG	23,00			chromosome 1 open reading frame 8 cytochrome c oxidase subunit IV
	GTACTGTAGCA	6,00			integrin, alpha 3 (antigen CD49C, alpha
.0.0		0,00	0,50	1 13.200023	3 subunit of
4514	TTCTTTCTCAC	1;00	0,14	Hs.9788	ESTs, Moderately similar to unnamed
		· 1			protein product
4515	CTCAGAGAACA	1,00	0,14	Hs.9414	KIAA1488 protein
	AAGCATCTCAG	1,00	0,14		transcobalamin II; macrocytic anemia
	TAGATCTGTAT	1,00	0,14	Hs.83419	KIAA0252 protein
	TACTTTATAAG	1,00	0,14	Hs.8230	a disintegrin-like and metalloprotease (reprolysin t
4519	CACAAATGCTG	1,00	0,14	Hs.80562	gelsolin (amyloidosis, Finnish type)
	CTGTGTCCAAG	1,00			epidermal growth factor receptor
					pathway substrate 1
	AAGAACTACAC	1,00			KIAA0172 protein
4522	GCAAGGAAAAA	1,00	0,14	Hs.77496	small nuclear ribonucleoprotein
					polypeptide G
	AAATGCAGTAG	1,00			cathepsin O
	GAAATTCAAAC	1,00		Hs.74649	cytochrome c oxidase subunit VIc
4525	ATGCCTTTGAA	1,00	0,14	Hs.7298	biphenyl hydrolase-like (serine
4520	TTCCCTTCCTT	4.00	0.44	II. 50004	hydrolase; breast ep
	TCCGTTCCTT	1,00			KIAA0867 protein
	TCGGTGTCTGG	1,00		Hs.4953	golgi autoantigen, golgin subfamily a, 3
4020	GTGGCTCAATT	1,00	U, 14	Hs.4864	KIAA0892 protein

4500	00770070	4 00	0.44	11- 05070	IZIA A OOFOto-i
	GGTTGGTGGTC	1,00			KIAA0852 protein
-	GGAGGTGCTCA	1,00			ESTs, Weakly similar to B9 [H.sapiens]
	GGCCTGGGGGT	1,00			vav 3 oncogene
-	TITGTTGTATG	1,00			ESTs
4533	GCATCTTCAAA	1,00			interferon-induced protein 41, 30kD
4534	GAGAAGCCCCG	1,00	0,14		ESTs, Weakly similar to ALU6_HUMAN ALU SUBFAMILY SP
4535	ACAGGCAGAAA	1,00	0,14	Hs.200526	TNF receptor-associated factor 2
4536	GCGTTCAATAA	1,00	0,14	Hs.19720	Homo sapiens mRNA full length insert cDNA clone EURO
4537	AAGCACTTCTG	1,00	0,14	Hs.189658	CGI-149 protein
4538	CAGTCCCGGCT	1,00	0,14	Hs.189658	CGI-149 protein
4539	TATATACATTT	1,00			peptidase (mitochondrial processing) beta
4540	TGACTGTCACG	1,00	0,14		Homo sapiens clone 23914 mRNA sequence
4541	GGCAGATTGCT	1,00	0,14	Hs.173259	uncharacterized bone marrow protein BM033
4542	TTCTCGAGATG	1,00	0,14	Hs.161554	hypothetical protein FLJ20159
4543	GGACCATTGAA	1,00	0,14	Hs.155244	pre-mRNA splicing factor similar to S. cerevisiae Pr
4544	CCTTTCTGTAA	1,00	0,14	Hs.143648	insulin receptor substrate 2
	TGGAAATCAAG	1,00		Hs.13405	
	AAGCCAGTCTG	1,00		Hs.112378	LIM and senescent cell antigen-like domains 1
4547	ACTITICAAAA	1,00	0,14		HSPC037 protein
4548	TATGTATTTCT	1,00			lysyl oxidase
4549	GTGCTATTCTG	4,00			Homo sapiens mRNA full length insert cDNA clone EURO
4550	TCTGTCCTCAG	4,00	0,42	Hs.75216	protein tyrosine phosphatase, receptor type, F
4551	GGGTGTGGTGG	4,00	0,42		Homo sapiens mRNA; cDNA DKFZp564N123 (from clone DKF
4552	GCCCCCAATAA	66,00			lectin, galactoside-binding, soluble, 1 (galectin 1)
4553	AAATATGAGCT	3,00	0,36	Hs.181368	U5 snRNP-specific protein (220 kD), ortholog of S. c
4554	GAGTCAGGAGA	9,00	0,59		CGI-120 protein
	GGGGCCCCCTC	4,00			Sjogren's syndrome nuclear autoantigen
4556	CTCACTTCTTA	4,00	0,42	Hs.165998	DKFZP564M2423 protein
	CTCTGCCCTCC	4,00			ESTs, Weakly similar to dJ68O2.2 [H.sapiens]
4558	СТСТСТСТСТТ	2,00	0,27	Hs.7833	selenium binding protein 1
	GACTATAGCGC	2,00		Hs.7036	N-Acetylglucosamine kinase
	TTTTGTGTATT	2,00			thioredoxin-like
	AAGGAAGATTG	2,00			hypothetical protein FLJ20343
	TGCTCAGTGGT	2,00			dynein, cytoplasmic, light intermediate polypeptide

4563	GCATTTAAATA	28,00	0,84	Hs.275959	eukaryotic translation elongation factor 1 beta 2
4564	CTGACCTGTGT	28,00		Hs.77961	major histocompatibility complex, class I, B
4565	AAACATTAGCC	3,00		Hs.82911	protein tyrosine phosphatase type IVA, member 2
	CTTTGTTTAAT	1,00			Homo sapiens mRNA; cDNA DKFZp434L235 (from clone DKF
	GAGAGGAGAGG	1,00	•	Hs.92002	guanine nucleotide binding protein (G protein), alph
4568	GAGAACTCCCC	1,00	0,14	Hs.82001	polycystic kidney disease 2 (autosomal dominant) -NO
4569	TAGTTGCAAAT	1,00			BTG family, member 3
4570	GATGCTAACCA	1,00	0,14	Hs.76591	KIAA0887 protein
4571	TACCTTTATTG	1,00	0,14	Hs.75761	SFRS protein kinase 1
4572	ATCACAGGTGA	1,00	0,14	Hs.7252	KIAA1224 protein
4573	AACAAGTCTTT	1,00	0,14	Hs.69851	GAR1 protein
4574	TGGGGGTTTCC	1,00	0,14	Hs.62954	ferritin, heavy polypeptide 1
4575	GCCAAGACACA	1,00	0,14		mel transforming oncogene (derived from cell line NK
4576	GAGACTGCAAT	1,00	0,14		solute carrier family 11 (proton-coupled divalent me
4577	TAAACAGGTGG	1,00	0,14	Hs.5473	ESTs, Moderately similar to ALU5_HUMAN ALU SUBFAMILY
4578	TCTGATCAGGA	1,00	0,14	Hs.50441	CGI-04 protein
4579	AAGAGACATAG	1,00		Hs.46847	TRAF and TNF receptor-associated protein
4580	AAGAAGCGCAA	1,00	0,14	Hs.46784	potassium large conductance calcium- activated channe
4581	GCACCCAACAC	1,00	0,14	Hs.4082	lectin, galactoside-binding, soluble, 8 (galectin 8)
4582	AGGGACTTGTG	1,00	0,14	Hs.29331	carnitine palmitoyltransferase I, muscle
4583	TTTGCAAAAAA	1,00	0,14	Hs.286083	
4584	TGTGTGTAA	1,00	0,14	Hs.272328	KRAB-zinc finger protein synten
4585	TTTGAGTTCTT	1,00	0,14	Hs.24789	ESTs
	AACTCTAAGGA	1,00		Hs.238797	
4587	TGGTACTTCTC	1,00	0,14	Hs.22353	ESTs, Weakly similar to AF151847_1 CGI-89 protein [H
4588	GGGTGAGGGGG	1,00	0,14	Hs.21840	Homo sapiens clone 24852 mRNA sequence
4589	GCTATTTCCTA	1,00	0,14	Hs.199009	Homo sapiens PCCX2 mRNA for protein containing CXXC
4590	GAGACGCATTT	1,00	0.14	Hs.184585	LIM domain only 2 (rhombotin-like 1)
	CTTGGTGCTGC	1,00			ribosomal protein L13
	ACCCGGTACAG	1,00			dihydropyrimidinase-like 2
	TATCCTCTGGC	1,00			Homo sapiens cDNA FLJ20290 fis, clone HEP04540
4594	AAACACCAAAT	1,00	0,14	Hs.146388	microtubule-associated protein 7
4595	CCCAGCCACTT	1,00			ribosomal protein S19

4506	COACTCCACTC	1 1 00	0.14	U- 42272	TCT- Woolds similarte ALLIA LILIMAN
	GCAGTGCACTC	1,00			ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J S
4597	GAAGGTTGTGG	1,00	0,14	Hs.12017	KIAA0439 protein; homolog of yeast ubiquitin-protein
4598	AAAGGCAGGAA	1,00	0,14	Hs.118152	
4599	TATCAGTGCCT	1,00			CGI-43 protein
4600	GCAAATGTACA	1,00	0,14	Hs.11238	KIAA0622 protein; Drosophila "multiple asters" (Mast
4601	CTTAGTCTAAA	1,00			damage-specific DNA binding protein 1 (127kD)
	ATGAAATAGTA	1,00			CED-6 protein
	GCTTGACATTG	1,00		Hs.103352	
	GTTCCCTGGCC	42,00			Finkel-Biskis-Reilly murine sarcoma virus (FBR-MuSV)
4605	CTGCAACCTAA	3,00	0,36	Hs.50785	SEC22, vesicle trafficking protein (S. cerevisiae)-l
	TGTGGTGGTGT	5,00			MLN51 protein
4607	GACAAAAAGTC	2,00	0,27	Hs.9683	protein-kinase, interferon-inducible double stranded
4608	ATCTATGACCC	2,00		Hs.8179	hypothetical protein, clone 2746033
4609	TAACCGCAGAT	2,00	0,27	Hs.74376	olfactomedin related ER localized protein
	CCTGCCGTCGG	2,00	0,27	Hs.130541	KIAA1542 protein
	ACCTCCCACCC	2,00			Human clone 23759 mRNA, partial cds
	GGGCCAATAAA	8,00			DKFZP566I1024 protein
	AAAAAACCCAA	8,00	0,57	Hs.111680	endosulfine alpha
	TTCAATAAAAA	93,00			ribosomal protein, large, P1
	TTCACTGCCGA	5,00			ATPase, vacuolar, 14 kD
	GCTTTTCAGAC	4,00			vascular endothelial growth factor B
	TAGTTGTAGGG	4,00			hypothetical protein
	TTTGGTGTTTG	4,00			F-box only protein 9
	TATCTTGCTTA	3,00			ring finger protein 11
	TATCTGGTCTT	3,00			signal transducer and activator of transcription 3 (
	GCTGGCCTG	9,00			H.sapiens clathrin light chain b gene
	TGTCTGTGGTA	5,00			HLA-B associated transcript-1
	AGGGCTTCCAA	104,00	1,21	Hs.29797	ribosomal protein L10
4624	TATAGGCCGAA	2,00	0,28		vitamin A responsive; cytoskeleton related
	ACTGTTCTCTT	2,00			Human clone 23722 mRNA sequence
4626	TCCTTTGTGCC	2,00	0,28		Homo sapiens HSPC283 mRNA, partial cds
	TTTATTGAATT	2,00	0,28	Hs.43910	CD164 antigen, sialomucin
	TATATGGATGT	2,00	0,28	Hs.42758	ESTs
	TCTAGTCACTG	2,00	0,28	Hs.36565	ESTs
4630	CAGCTCTTAGG	2,00	0,28	Hs.22208	Homo sapiens mRNA full length insert cDNA clone EURO
4631	GGACCTTGGAG	3,00	0,36	Hs.78877	inositol 1,4,5-trisphosphate 3-kinase B
	TATATTGATTG	3,00			B-cell translocation gene 1, anti- proliferative

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4633	ATTACAAACCT	3,00	0.36	Hs.30376	hypothetical protein
	CAGTGGGGTTA	3,00			hypothetical protein FLJ20303
	CTCCTCACCTG	47,00			lecithin-cholesterol acyltransferase
-	TACATTGCTTT	4,00		Hs.75104	RNA-binding protein S1, serine-rich
		.,,	-,		domain
4637	CTGGCTGCAAA	14,00	0,70	Hs.1342	cytochrome c oxidase subunit Vb
	GGTGGGGAGAT	7,00	0,54	Hs.157236	membrane protein of cholinergic synaptic
					vesicles
4639	AAGCCTTGCTG	6,00		Hs.6289	growth factor receptor-bound protein 2
	TTAGGGAGGAG	1,00		Hs.99995	intercellular adhesion molecule 3
	GAAAAGCTCCT	1,00		Hs.99843	DKFZP586N0721 protein
4642	GAAGGTCCTGC	1,00	0,14	Hs.979	pyruvate dehydrogenase (lipoamide) beta
4643	GATCTGAGGAG	1,00	0,14	Hs.96984	Homo sapiens cDNA FLJ20335 fis, clone
					HEP11429 '
	ATGAAAAGTGC	1,00	0,14	Hs.78743	zinc finger protein 131 (clone pHZ-10)
	ATATGAAGCAT	1,00		Hs.7862	hypothetical protein FLJ20312
4646	GGAGGGACCCC	1,00	0,14	Hs.78473	N-deacetylase/N-sulfotransferase
				<del></del>	(heparan glucosamin
	GGAGACTTCCT	1,00			annexin A4
4648	GCTCTGCCCTC	1,00	0,14	Hs.68257	general transcription factor IIF,
10.10		4.00		11 00001	polypeptide 1 (74k
	AGGTGTCTTTG	1,00			KIAA0937 protein
4650	AACAAATCCTT	1,00	0,14	Hs.5169	suppressor of G2 allele of SKP1, S.
4651	CAGGCGTGCAC	1,00	0.14	Hs.33818	cerevisiae, homo RecQ protein-like 5
	TATAAATTAAA	1,00			thyroid receptor interacting protein 15
	ACACACAAAA	1,00			KIAA1007 protein
	AAGCAAGAATG	1,00			KIAA1376 protein
	ACTTAAGGTTG	1,00			RAB10, member RAS oncogene family
	GTGTCCTTGTC	1,00			Homo sapiens mRNA; cDNA
	0.0.000.0	.,00	0,		DKFZp566H2446 (from clone DK
4657	CTCTCCTGCTC	1,00	0.14		PR domain containing 4
	GCCAAGTGAAC	1,00	0,14	Hs.20225	tuftelin-interacting protein
4659	AGGCCCCAGGG	1,00			putative Rab5 GDP/GTP exchange
					factor homologue
	TAACAAGTTTC	1,00	0,14	Hs.179902	Homo sapiens CTL1 gene
4661	GTAGTCCAAAC	1,00	0,14	Hs.178617	ESTs, Weakly similar to AF151840_1 CGI-82 protein [H
4662	GTGATGTCTGT	1,00	0,14	Hs.164026	
4663	GTCTACAATTG	1,00			hypothetical protein FLJ20159
4664	ATAATTGACTA	1,00			COP9 subunit 6 (MOV34 homolog, 34 kD)
4665	TTGGTGAAGGA	70,00	1.12	Hs.75968	thymosin, beta 4, X chromosome
	GCTTGTTAAGA	3,00		Hs.4113	S-adenosylhomocysteine hydrolase-like
		=,=9	5,50		1
4667	ATTGTGCTTGC	3,00	0,36	Hs.251531	proteasome (prosome, macropain) subunit, alpha type,
4668	AGGAAAAAAA	3,00	0.36	Hs.180639	
			- 3,00		

-	·				
	GCTGGGAGGG	4,00		Hs.20733	
	AGTGTGCGCTT	2,00			Homo sapiens GT212 mRNA
	AACTTGGCTGT	2,00		Hs.79107	
	GGGGACTGGTG	2,00			protein kinase C, delta
4673	CCAGCTGCCAA	10,00	0,63	Hs.2055	ubiquitin-activating enzyme E1 (A1S9T
					and BN75 tempe
	AATAGGTCCAA	57,00			ribosomal protein S25
4675	TTGGCATTGTC	3,00	0,36	Hs.250911	Homo sapiens clone 23967 unknown
					mRNA, partial cds
	CCCCAGTTGCT	36,00		Hs.74451	calpain 4, small subunit (30K)
4677	GCATAATAGGT	90,00	1,26	Hs.184108	ribosomal protein L21 (gene or pseudogene)
4678	CTCTGATGCAG	2,00		Hs.80961	polymerase (DNA directed), gamma
4679	GTCATTATGCT	2,00	0,28	Hs.78305	RAB2, member RAS oncogene family
	TAAACTGTTAA	2,00	0,28	Hs.3491	ribosomal protein S14
4681	CTAGCCAGCAG	2,00	0,28	Hs.24983	hypothetical protein from EUROIMAGE
					2021883
	GCTAAACTCTG	2,00		Hs.207409	
	AGACGCTTCTG	2,00	0,28	Hs.203772	FSHD region gene 1
4684	GATCTGTTTCT	2,00	0,28	Hs.169743	Homo sapiens clone 25121 neuronal
1000					olfactomedin relat
	CTGGCCTGTGT	2,00			villin 2 (ezrin)
4686	GCTGTTTAAAA	2,00	0,28	Hs.154320	ubiquitin-activating enzyme E1C
4697	CCCTCTAATAA		0.47	11- 0540	(homologous to yeast
	CCCTGTAATAA	5,00			selenoprotein N
	TCTGGGGACGA ATTTCTTGCCG	5,00	0,47	HS.74111	RNA-binding protein (autoantigenic)
4009	ATTICITIECCE	5,00	0,47	Hs.25682	Homo sapiens HSPC262 mRNA, partial cds
4690	ATTAAATTCAG	3,00	0.36	He 218320	hypothetical protein
	CCTTTCAAGCA	3,00			eukaryotic translation initiation factor 3,
		0,00	0,00	113.130033	subunit
	TAATCCCAGCA	4,00		Hs.236710	EST
	TCTTCCCCAGT	4,00	0,42	Hs.14231	selenoprotein W, 1
4694	TGCCTGCACCA	41,00	1,00	Hs.135084	cystatin C (amyloid angiopathy and
					cerebral hemorrha
	CCGTGGTCGTG	10,00	0,63		fibrillarin
4696	GGGAAACCCTG	5,00	0,47	Hs.161137	ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB
	GCTTCTGCATA	1,00	0,14	Hs.8886	hypothetical protein FLJ20424
4698	GTTGCATTTTC	1,00	0,14	Hs.81001	F-box only protein 25
	GTGATACGTTG	1,00	0,14	Hs.80306	Homo sapiens mRNA, clone:RES4-4
4700	CAGTTTGAAAT	1,00	0,14	Hs.77256	enhancer of zeste (Drosophila) homolog 2
4701	CAGTGATTCCA	1,00	0,14	Hs.75056	adaptor-related protein complex 3, delta 1 subunit
4702	CATATTTGTGA	1,00	0.14	Hs.6489	dynactin p62 subunit
	TACATTTGCAA	1,00		Hs.6349	Human Chromosome 16 BAC clone CIT987SK-A-362G6
4704	TAATTGCAGAT	1,00	0 14	Hs.4854	cyclin-dependent kinase inhibitor 2C
		•,00		1 10: 1007	1979mi dopondoni kindoe ininbitoi 20

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	T			<u> </u>	(p18, inhibits
4705	CCATCCGCAGG	1,00	0.14	Hs.4437	ribosomal protein L28
					<del></del>
	CAATGGAGCTT	1,00		Hs.30925	hypothetical protein FLJ10199
_	TTGTACCACCT	1,00		Hs.29024	ESTs
	GCCTCCAGATT	1,00		Hs.278355	
	TAAAATAAAAG	1,00			hypothetical protein from EUROIMAGE 363668
4710	GTATGGACTAT	1,00	0,14	Hs.268012	fatty-acid-Coenzyme A ligase, long-chain 3
4711	AACCTGTTCTA	1,00	0,14	Hs.26761	DKFZP586L0724 protein
4712	TGTGGCCCACC	1,00	0,14	Hs.250870	mitogen-activated protein kinase kinase 5
4713	TTTGTGGTCAA	1,00	0,14	Hs.244099	EST, Weakly similar to EF2_HUMAN ELONGATION FACTOR 2
4714	ACATTITTAAC	1,00	0,14	Hs.177516	high density lipoprotein binding protein (vigilin)
	TGCCCCGCACT	1,00	0,14	Hs.169119	
4716	CCTGGGGGCCG	1,00			nuclear transcription factor Y, gamma
4717	GATTTAAATCA	1,00			transcriptional intermediary factor 1 gamma
4718	AGCCCTGGCTG	1,00	0.14	Hs.15896	
	GGCGCACTCTG	1,00			Homo sapiens clone 24706 mRNA
		,,,,,	٠,٠٠		sequence
4720	CAGGAGAACTG	1,00	0,14	Hs.150614	ESTs, Weakly similar to ALU4_HUMAN ALU SUBFAMILY SB2
4721	AAAACTTTTGT	1,00	0,14	Hs.126219	hypothetical protein FLJ20081
4722	AACTCTCAATG	1,00	0,14	Hs.12150	retinal short-chain
					dehydrogenase/reductase retSDR2
_	AGGAAACTGGG	1,00		Hs.11955	
4724	TTTTGCAACAA	1,00	0,14	Hs.118739	ESTs, Weakly similar to JC2473 doc2 protein - human
4725	GGAAAATTGGT	1,00	0,14	Hs.115740	KIAA0210 gene product
4726	GAACTGTGAGT	1,00		Hs.110044	
4727	TACACCAGCAA	1,00			ESTs, Weakly similar to Homolog of rat Zymogen granu
4728	TGCAGCACGAG	18,00	0,78	Hs.110309	major histocompatibility complex, class I,
4729	CTGGATGGGCA	6,00	0,52	Hs.44017	SIR2 (silent mating type information regulation 2, S
4730	CAGGGAGCGCC	3,00	0,36	Hs.8657	TPA inducible gene-1
	CGTGTGCCTGT	3,00		Hs.74649	cytochrome c oxidase subunit VIc
	TTCTCTCCACA	2,00		Hs.94446	polyamine-modulated factor 1
	CCCCATACTAC	2,00		Hs.57652	EGF-like-domain, multiple 2
	TTGTAATAAAA	2,00			SMC (mouse) homolog, X chromosome
	TTTACAAATAA	2,00			KIAA0493 protein
	TTAACCCTCTA	8,00			H3 histone, family 3B (H3.3B)
	AAGGTAGCAGA	8,00			adenylyl cyclase-associated protein
	GACCTCCTGCC	3,00		Hs.89449	mitogen-activated protein kinase kinase kinase 11

4739	TTACCATATCA	39,00	1.02	Hs 177461	ribosomal protein L39
	AAATACAGCAG	4,00	0.43	Hs.182429	protein disulfide isomerase-related
		.,55	•,		protein
4741	CCCGTCCGGAA	145,00	1,66	Hs.180842	ribosomal protein L13
	TTGAGCCAGCC	11,00			KH-type splicing regulatory protein
					(FUSE binding pr
4743	CCTGAGGTCAG	2,00	0,28	Hs.95196	ESTs, Weakly similar to T20B12.3
	,				[C.elegans]
	TAACATTGGTG	2,00		Hs.79306	eukaryotic translation initiation factor 4E
	GTAGAAAAAA	2,00			numb (Drosophila) homolog
	GGGCTCTGAGC	2,00			LCAT-like lysophospholipase
	CTTTTCATCAT	2,00			x 003 protein
	TCACTGATCTT	2,00		Hs.3491	ribosomal protein S14
	GCTGGAATAAA	2,00			hypothetical protein FLJ20211
	TTGGCCGGGCT	2,00		Hs.248488	
4751	AAAGTCTAGAA	15,00	0,75	Hs.82932	cyclin D1 (PRAD1: parathyroid
1					adenomatosis 1)
4752	GAGCTTTTGAA	3,00	0,37		Homo sapiens cDNA FLJ11066 fis, clone PLACE1004885
4753	TCCTAGCCTGT	4,00	0,43		splicing factor similar to dnaJ
4754	CCTGAAATTTG	5,00	0,48		heterogeneous nuclear ribonucleoprotein
					A0
4755	TTAATCCTAAA	5,00	0,48	Hs.150741	2',3'-cyclic nucleotide 3'
					phosphodiesterase
	ATAATTCTTTG	93,00		Hs.539	ribosomal protein S29
	ACATTTCATTT	1,00		Hs.8966	integral membrane protein 1
4758	TCCAGGGCCĞC	1,00	0,14		ESTs, Weakly similar to S57447 HPBRII-
4750	GAACCACAGGA	1.00	0.14	Hs.80042	7 protein - hu dolichyl-P-Glc:Man9GlcNAc2-PP-
4/09	GAACCACAGGA	1,00	0, 14		dolichylglucosyltransf
4760	AATATTTAGTG	1,00	0 14		transcription factor 6-like 1 (mitochondrial
7700		1,00	0, 14		transcr
4761	ACTITITGGCT	1,00	0.14	Hs.6968	KIAA1460 protein
-	GAAGCAAAAA	1,00		Hs.6278	DKFZP586B1621 protein
4763	GCCGCTGCCAG	1,00		Hs.6162	KIAA0771 protein
4764	CTCAGAACACT	1,00		Hs.5309	ESTs
4765	ATGGTCTCCTG	1,00			ESTs
4766	TITGTTTTAT	1,00	0,14	Hs.3622	procollagen-proline, 2-oxoglutarate 4-
					dioxygenase (p
	AGTACCTTATA	1,00			PRO0529 protein
	CCTGAAAAAA	1,00			PRO1578 protein
	AATGCCCCACT	1,00			hypothetical protein FLJ10830
4770	TCAGACTAGTT	1,00	0,14	Hs.268012	fatty-acid-Coenzyme A ligase, long-chain
4771	TCTGGCAAAGA	1,00	0.14	He 267821	Rho GTPase activating protein 5
	AGGCGAGCTGC	1,00			hypothetical protein PRO2577
	GTTCATTTTGA	1,00			hypothetical protein FKO2377
	TTAAAGAGCCG	1,00			growth arrest-specific 7
	GAATTTCCCAG	1,00		Hs.2253	complement component 2
	10.000AG	1,00		1. 10.2200	Positions combonent &

<u></u>		4.5-	6.1	1.1 46	
<u> </u>	AAAAATAAAGA	1,00			ESTs, Weakly similar to INI7_HUMAN INTERFERON-ALPHA
4777	GCAACTTTTTG	1,00	0,14	Hs.183994	protein phosphatase 1, catalytic subunit, alpha isof
4778	AGCGGCTACAC	1,00	0,14	Hs.183487	interferon stimulated gene (20kD)
4779	TGCAACTACAA	1,00	0,14	Hs.182278	calmodulin 2 (phosphorylase kinase, delta)
4780	GGCCTGGAATT	1,00	0,14	Hs.171566	hypothetical protein
4781	TATTCCTGTGA	1,00			karyopherin (importin) beta 2
4782	ATGCCCAATGT	1,00	0,14	Hs.155396	nuclear factor (erythroid-derived 2)-like 2
4783	GAAATAAAAAG	1,00	0,14	Hs.154156	myosin, light polypeptide 4, alkali; atrial, embryon
4784	GCCTTTCTAAT	1,00	0,14	Hs.149957	ribosomal protein S6 kinase, 90kD, polypeptide 1
4785	CATTTTACTGG	1,00	0,14	Hs.147189	HYA22 protein
	GCCAGATTGAG	1,00			CGI-145 protein
4787	TTTTATTAAAG	1,00	0,14	Hs.108504	hypothetical protein FLJ20113
4788	AAGTGGAATAA	1,00			KIAA0404 protein
4789	GATCTCACTGT	1,00			KIAA0477 gene product
4790	GAGACTCCTGC	10,00	0,65		solute carrier family 2 (facilitated glucose transpo
4791	TGCACTTCAAG	13,00	0,71		SPARC-like 1 (mast9, hevin)
4792	AAAGAAAGTGG	5,00			mannosyl (alpha-1,3-)-glycoprotein beta- 1,2-N-acetyl
4793	TCCATCTGTTG	4,00	0,43	Hs.252189	syndecan 4 (amphiglycan, ryudocan)
4794	ATCCACCCGCC	3,00	0,37	Hs.251337	ESTs
	CTGCTTCCTGA	2,00			A kinase (PRKA) anchor protein 1
4796	TGCTTATTGAA	2,00		Hs.5822	lectin, mannose-binding, 1
4797	ACTCACGATTG	2,00	0,28	Hs.4814	mannosidase, alpha, class 1B, member 1
4798	GGATGTGGAGG	2,00	0,28	Hs.31305	transducin-like enhancer of split 3, homolog of Dros
4799	AATTCCCGTCC	2,00	0,28	Hs.18349	HSPC145 protein
4800	AGCACATTTGA	8,00	0,59		gelsolin (amyloidosis, Finnish type)
4801	CACTACACGGG	6,00	0,53	Hs.227729	FK506-binding protein 2 (13kD)
	GTTCAAAGACT	3,00	0,37	Hs.75260	mitogen inducible 2
4803	CTCAAGCACCA	3,00	0,37	Hs.279652	CGI-28 protein
4804	ACCACTTATCC	3,00	0,37	Hs.249982	cathepsin B
	GACATAAATCC	5,00			Nef-associated factor 1
4806	GGTTCCTGGTG	2,00	0,28	Hs.3074	ESTs, Weakly similar to KIAA0386 [H.sapiens]
4807	ATTTCTGCTGG	2,00	0,28	Hs.279607	calpastatin
	GCTCTGTAAGC	2,00			putative methyltransferase
4809	GTGAGACCCCC	2,00		Hs.235989	
4810	CTGTTAATAAA	2,00	0,28	Hs.19500	nuclear localization signal deleted in velocardiofac
4811	GTTGAGTAACA	2,00	0,28	Hs.184776	ribosomal protein L23a
	CAACTTTAGGG	2,00			heterogeneous nuclear ribonucleoprotein D-like

4813	GCTACTATTAG	2,00	0.28	He 154443	minichromosome maintenance deficient
					(S. cerevisiae)
4814	ATGTGAAGAAT	2,00	0,28	Hs.13662	Homo sapiens clone 25036 mRNA sequence
4815	ACCTGCCCCTC	2,00	0,28	Hs.125262	DKFZP586G1624 protein
4816	CCTGTCCTGCA	7,00	0,57	Hs.11417	Rab acceptor 1 (prenylated)
4817	CCTTTGTAAGT	5,00			v-jun avian sarcoma virus 17 oncogene homolog
4818	AAGGACCTTTT	12,00	0,71	Hs.109051	glycoprotein, synaptic 2
4819	ATCGGGCCCGG	10,00	0,66	Hs.274411	SCAN domain-containing 1
4820	AAAAATAAAGC	3,00			WD repeat domain 1
4821	ACTGATCTGCA	3,00	0,37	Hs.264190	vacuolar protein sorting 35 (yeast homolog)
4822	TCTCTGCAAAA	3,00	0,37	Hs.25489	hypothetical protein FLJ20640
4823	AGCCTGGACTG	7,00			cell membrane glycoprotein, 110000M(r) (surface anti
4824	CCCCGCCAAGT	8,00	0,60	Hs.169718	calponin 2
4825	TCAAAAAAAA	11,00			deleted in lung and esophageal cancer 1
	TTATAACTGAA	7,00	0,57	Hs.79933	cyclin I
4827	TGCTGCCTCAG	2,00	0,28	Hs.30792	hook2 protein
4828	CTAGTCACTTC	2,00	0,28	Hs.180577	granulin
4829	AATGAGCAACT	2,00	0,28		guanylate binding protein 2, interferon- inducible
4830	CAATTCCTTCA	2,00	0,28	Hs.170098	KIAA0372 gene product
4831	AGCTGGGATGG	2,00			peroxisomal 2,4-dienoyl-CoA reductase
4832	CCACGTGGCTG	2,00			semaphorin Rs, short form
4833	GCACCTTCTGG	2,00	0,28	Hs.132744	hypothetical protein
4834	CCCTGTTTTT	1,00	0,15	Hs.92030	ESTs
4835	TTTTTCTTCA	1,00	0,15	Hs.8262	lysosomal-associated membrane protein 2
4836	TTCTAGACCAT	1,00	0,15	Hs.78946	cullin 3
4837	GAGATGAAATG	1,00	0,15	Hs.75939	uridine monophosphate kinase
4838	GTCTGTGTATG	1,00		Hs.7481	ESTs, Weakly similar to similar to mitochondrial RNA
4839	CCAAGAGGAAA	1,00	0,15	Hs.74649	cytochrome c oxidase subunit VIc
4840	AATCAAGGTGT	1,00	0,15	Hs.73287	KIAA1235 protein
4841	CCCTGAGGCCA	1,00	0,15	Hs.6607	Homo sapiens mRNA; cDNA DKFZp566F164 (from clone DKF
4842	TCTGAAGACTT	1,00	0,15	Hs.5548	f-box and leucine-rich repeat protein 5
4843	TTGGATATGTG	1,00	0,15	Hs.554	Sjogren syndrome antigen A2 (60kD, ribonucleoprotein
4844	CATTGTCTTCA	1,00	0,15	Hs.5054	CGI-133 protein
4845	TGTGAAGATTA	1,00	0,15	Hs.44829	ESTs
4846	AAGTGAAGAGC	1,00		Hs.31945	hypothetical protein FLJ11071
4847	TCGTCCTAGAA	1,00			KIAA0685 gene product
4848	ACTGATAACAG	1,00		Hs.27182	phospholipase A2-activating protein
	ATTTTGGATTC	1,00			ESTs
	CAGTGCCCAAC	1,00		Hs.26409	ESTs
	CATTGGTAGAA	1,00			Homo sapiens clone 23967 unknown

	<u></u>	r - (		T	mRNA, partial cds
4852	TGAAAATCAAA	1,00	0.15	Hs 239489	TIA1 cytotoxic granule-associated RNA-
1002	10,000110,001	',55	0,10	13.200400	binding protei
4853	AGGCTGCGGTG	1,00	0.15	Hs 209646	KIAA1118 protein
	TAACTCCATTG	1,00			SEC24 (S. cerevisiae) related gene
-00-	1741010071110	',55	0,10	113.10022	family, member D
4855	CCGCCCTTCGG	1,00	0.15	Hs.180677	zinc finger protein 162
	ACTTAACATTT	1,00			heat shock 90kD protein 1, alpha
	AGGCTTCTAGC	1,00			Homo sapiens mRNA; cDNA
		.,	0,.0		DKFZp434E1515 (from clone DK
4858	GTGACTGCCAG	1,00	0,15	Hs.177536	metallocarboxypeptidase CPX-1
	ATTAGTCAGAA	1,00			inhibin, beta B (activin AB beta
					polypeptide)
4860	TTAACACTGTG	1,00	0,15	Hs.172069	DKFZP434C212 protein
4861	AATTACCAAAG	1,00	0,15	Hs.170218	KIAA0251 protein
4862	GTATCTTAATA	1,00	0,15	Hs.155489	NS1-associated protein 1
4863	TGTTCTCAAGT	1,00		Hs.154396	
4864	CCTAAGGGAGA	1,00	0,15	Hs.153022	TATA box binding protein (TBP)-
					associated factor, RN
4865	TCTCCACGAAG	1,00	0,15	Hs.149570	actin related protein 2/3 complex, subunit
					4 (20 kD)
	GTTTGGATCTG	1,00		Hs.118836	
	TAAGATTAGAA	1,00			timeless (Drosophila) homolog
4868	TATTTGCTACA	1,00	0,15		glucocorticoid receptor DNA binding
4000	07004700407	2.22			factor 1
4869	GTGGATGGACT	3,00	0,37		seven transmembrane domain orphan
1970	CCCAATAAACC	2.00	0.27	H= 2020	receptor
	GGGAATAAACC ATCAGTGTGCA	3,00 3,00		Hs.3828	mevalonate (diphospho) decarboxylase calponin 3, acidic
	GCCATAAAATG	3,00		Hs.1908	
	AAGTTGCTATT	23,00		Hs.78575	proteoglycan 1, secretory granule prosaposin (variant Gaucher disease and
40/3	ANGIIGCIAII	23,00	0,92	па. горго	variant meta
4874	CTCATAGCAGT	14,00	0.76	Hs 119252	tumor protein, translationally-controlled 1
	GGCTGAGAATG	6,00	0.54	Hs 106845	ESTs, Weakly similar to ALU6_HUMAN
	00010/10/110	0,00	0,04	110.100040	ALU SUBFAMILY SP
4876	GGATGTGAAAG	9,00	0.64	Hs.177543	antigen identified by monoclonal
		3,33			antibodies 12E7, F2
4877	TTACACCTGTG	2,00	0,29	Hs.82794	centrin, EF-hand protein, 2
	CCACCTGCTTT	2,00	0,29	Hs.7367	Homo sapiens glucocorticoid receptor
					AF-1 coactivato
4879	AGCACTGTACT	2,00	0,29	Hs.6375	uncharacterized hypothalamus protein
					HT010
	CCCGGCTCCTC	2,00			hypothetical protein FLJ20041
	GTTTTCCTGAA	2,00			ESTs
4882	TGCCTCCCATC	2,00	0,29	Hs.2437	eukaryotic translation initiation factor 2B,
1000	07010	ابيا			subunit
4883	GTGACGTGCAC	2,00	0,29	Hs.207390	ESTs, Moderately similar to
4004	COTTOOCTOO	0.55		11 4==0.55	ALU5_HUMAN ALU SUBFAMILY
4884	GCTTGGCTCCC	2,00	0,29	Hs.175260	IF21

4886 CTTTTTGTGCC         8,00         0,81 Hs.182328 GW/28 protein           4887 GAAGCAGGACC         67,00         1,46 Hs.180370 cofilin 1 (non-muscle)           4888 TGGAGTAGTAA         4,00         0,44 Hs.8975 PRO1073 protein           4889 AAGCGGGACCT         4,00         0,44 Hs.153436 Nacetyltransferase, homolog of S. cerevisiae ARD1           4890 GTTTCTATCAA         4,00         0,44 Hs.12540 Nysophospholipase I yophospholipase I yophospholipase I yophospholipase I yophospholipase I yophospholipase I yophospholipase I activator subunit 1           4892 ACAAAATAAAA         3,00         0,38 Hs.83469 nuclear factor (erythroid-derived 2)-like 1 nuclear factor (erythroid	4885	GGGGCAGGTC	2,00	0.29	Hs 107374	chromobox homolog 6
4887 [GAAGCAGGACC         67,00         1,46   Hs. 180370   PRO1073 protein           4888 [TTGTGATGTAA]         4,00         0,44   Hs. 1853436   PRO1073 protein           4889 [AGCGGGACCT]         4,00         0,44   Hs. 153436   N-acetyltransferase, homolog of S. cerevisiae ARD1           4890 [GTTCTATCAA]         4,00         0,62   Hs. 75348   proteasome (prosome, macropain) activator subunit 1           4892 [ACAAAATAAAA]         3,00         0,38   Hs. 83469   nuclear factor (erythroid-derived 2)-like 1           4893 [CATTGCAGGAT]         3,00         0,38   Hs. 83469   nuclear factor (erythroid-derived 2)-like 1           4894 [GTCCCTCTCAA]         2,00         0,29   Hs. 92198   calcium-regulated heat-stable protein (24kD)           4895 [TGAACAGTAAT]         2,00         0,29   Hs. 80120   UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-ace           4896 [TCCAGAATCCT]         2,00         0,29   Hs. 75243   bromodomain-containing 2           4897 [AACAGTGGCT]         2,00         0,29   Hs. 179565   minichromosome maintenance deficient (S. cerevisiae)           4899 [CTGCCTCCGTG]         2,00         0,29   Hs. 107014   membrane interacting protein of RGS16           4900 [ATTACACACG]         2,00         0,29   Hs. 107014   membrane interacting protein A8 (calgranulin A)           4901 [TACCTGCAGAA]         1,00         0,15   Hs. 99519   ESTs           4902 [CAAATGATGA]         1,00         0,15						
4888 TTGTGATGTAA         4,00         0,44 Hs.6975         PRO1073 protein           4889 AAGCGGACCT         4,00         0,44 Hs.153436         N-acetyltransferase, homolog of S. cerevisiae ARD1           4890 GTTTCTATCAA         4,00         0,44 Hs.12540         lysophospholipase I           4891 GAGATCCGCAA         8,00         0,62 Hs.75348         proteasome (prosome, macropain) activator subunit 1           4892 ACAAAATAAAA         3,00         0,38 Hs.83499         nuclear factor (erythroid-derived 2)-like 1           4893 CATTGCAGGAT         2,00         0,29 Hs.82198         calcium-regulated heat-stable protein (24kD)           4895 TGAACAGTAAT         2,00         0,29 Hs.7644         KIAA0469 gene product           4896 TCCAGAATCCT         2,00         0,29 Hs.7764         KIAA0469 gene product           4897 AACAACTGGCT         2,00         0,29 Hs.7764         KIAA0469 gene product           4899 CTGCCTCCGTG         2,00         0,29 Hs.179565 minichromosome maintenance deficient (S. cerevisiae)           4899 CTGCCTCCGTG         2,00         0,29 Hs.107014 membrane interacting protein of RGS16           4890 TACCGCGCTCC         1,00         0,15 Hs.99519         ESTs           4901 TACCTGCAGAA         1,00         0,15 Hs.89519         ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.8						
4889 AAGCGGGACCT         4,00         0,44 Hs.153436 N-acetyltransferase, homolog of S. cerevisiae ARD1           4890 GTTTCTATCAA         4,00         0,44 Hs.12540 Iysophospholipase I Iysophosphosphosphosphosphosphosphosphosph						
cerevisiae ARD1						
AB91   GAGATCCGCAA   8,00   0,62   Hs.75348   proteasome (prosome, macropain)   activator subunit   4892   ACAAAATAAAA   3,00   0,38   Hs.83469   nuclear factor (erythroid-derived 2)-like 1   4893   CATTGCAGGAT   3,00   0,38   Hs.4288   hypothetical protein DKFZp434K046   4894   GTCCCTCTCAA   2,00   0,29   Hs.92198   calcium-regulated heat-stable protein (24kD)   UIDP-N-acetyl-alpha-D-galactosamine:polypeptide N-ace   4895   TGAACAGTAAT   2,00   0,29   Hs.7764   KIAA0469 gene product   4897   AACAACTGGCT   2,00   0,29   Hs.7764   KIAA0469 gene product   4897   AACAACTGGCT   2,00   0,29   Hs.179565   minichromosome maintenance deficient (s. cerevisiae)   4899   CTGCCTCCGTG   2,00   0,29   Hs.179565   minichromosome maintenance deficient (s. cerevisiae)   4890   ATTACACCACG   2,00   0,29   Hs.107014   membrane interacting protein of RGS16   4901   TACCTGCAGAA   2,00   0,29   Hs.107014   membrane interacting protein of RGS16   4901   TACCTGCAGAA   2,00   0,15   Hs.99519   ESTs   4904   CTGGGATGCAG   1,00   0,15   Hs.99321   ESTs   4904   CTGGGATGCAG   1,00   0,15   Hs.93231   ESTs   4904   CTGGGATGCAG   1,00   0,15   Hs.89434   drebrin 1   4905   CAACTATCCGT   1,00   0,15   Hs.89434   drebrin 1   4905   CAACTATCCGT   1,00   0,15   Hs.89434   drebrin 1   4908   GCTTGACAGA   1,00   0,15   Hs.89434   drebrin 1   4908   GCTTGACAGA   1,00   0,15   Hs.6272   retinoblastoma-binding protein 2   4907   ATACACTITGT   1,00   0,15   Hs.6272   retinoblastoma-binding protein 2   4909   GCGCAGAGAG   1,00   0,15   Hs.6278   DKFZP586B1621 protein   4911   TGAAGTCAAA   1,00   0,15   Hs.6278   DKFZP586B1621 protein   4911   TGAAGTCAAA   1,00   0,15   Hs.265960   hypothetical protein FLJ10563   4914   GAAATCCCAAT   1,00   0,15   Hs.265960   hypothetical protein   4916   GAGGAGAGAAA   1,00   0,15   Hs.265960   hypothetical protein   4916   GAGGAGAGAA   1,00   0,15   Hs.269560   hypothetical protein   4916   GAGGAGAGAA   1,00   0,15   Hs.269560   hypothetical protein   4916   GAGGAGGTGGA   1,00   0,15   Hs.269560   hypothetical prote						cerevisiae ARD1
activator subunit 1						
4893 CATTGCAGGAT         3,00         0,38 Hs.4288         hypothetical protein DKFZp434K046           4894 GTCCCTCTCAA         2,00         0,29 Hs.92198         calcium-regulated heat-stable protein (24kD)           4895 TGAACAGTAAT         2,00         0,29 Hs.80120         UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-ace           4896 TCCAGAATCCT         2,00         0,29 Hs.75243         bromodomain-containing 2           4898 CAGGTCAAGAA         2,00         0,29 Hs.12851         minichromosome maintenance deficient (S. cerevisiae)           4899 CTGCCTCCGTG         2,00         0,29 Hs.12851         ESTs, Weakly similar to PSS1_HUMAN PHOSPHATIDYLSERIN           4900 ATTACACCACG         2,00         0,29 Hs.107014 membrane interacting protein of RGS16           4901 TACCTGCAGAA         2,00         0,29 Hs.100000 S100 calcium-binding protein A8 (calgranulin A)           4902 CCAAATGATGA         1,00         0,15 Hs.99519         ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.89321         ESTs           4905 CAACTATCCGT         1,00         0,15 Hs.89595         translocase of inner mitochondrial membrane 22 (yeas           4906 GCTTCATCAGA         1,00         0,15 Hs.75193         COP9 homolog           4907 ATACACTTTGT         1,00         0,15 Hs.6322         hypothetical protein FLJ10913           490	4891	GAGATCCGCAA	8,00	0,62	Hs.75348	
4894 GTCCCTCTCAA         2,00         0,29 Hs.92198         calcium-regulated heat-stable protein (24kD)           4895 TGAACAGTAAT         2,00         0,29 Hs.80120         UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-ace           4896 TCCAGAATCCT         2,00         0,29 Hs.775243         kIAAO469 gene product           4897 AACAACTGGCT         2,00         0,29 Hs.79565 minichromosome maintenance deficient (S. cerevisiae)           4898 CAGGTCAAGAA         2,00         0,29 Hs.19565 minichromosome maintenance deficient (S. cerevisiae)           4890 CTGCCTCCGTG         2,00         0,29 Hs.107014 membrane interacting protein of RGS16           4901 TACACACACG         2,00         0,29 Hs.100000         S100 calcium-binding protein A8 (calgraudin A)           4902 CCAAATGATGA         1,00         0,15 Hs.99519         ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.99321         ESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.93231         ESTs           4905 CAACTATCCGT         1,00         0,15 Hs.89434         drebrin 1         translocase of inner mitochondrial membrane 22 (yeas           4908 GCTTCATCAGA         1,00         0,15 Hs.76272         retinoblastoma-binding protein 2           4908 GCTTGGAGTTA         1,00         0,15 Hs.6336         upurping translocation breakpoint           4	4892	ACAAAATAAAA	3,00	0,38	Hs.83469	nuclear factor (erythroid-derived 2)-like 1
(24kD)	4893	CATTGCAGGAT	3,00	0,38	Hs.4288	hypothetical protein DKFZp434K046
Galactosamine:polypeptide N-ace	4894	GTCCCTCTCAA	2,00	0,29		(24kD)
4896 TCCAGAATCCT         2,00         0,29 Hs.7764         KIAA0469 gene product           4897 AACAACTGGCT         2,00         0,29 Hs.75243         bromodomain-containing 2           4898 CAGGTCAAGAA         2,00         0,29 Hs.179565         minichromosome maintenance deficient (S. cerevisiae)           4899 CTGCCTCCGTG         2,00         0,29 Hs.12851         ESTs, Weakly similar to PSS1_HUMAN PHOSPHATIDYLSERIN           4900 ATTACACCACG         2,00         0,29 Hs.100000         S100 calcium-binding protein of RGS16           4901 TACCTGCAGAA         2,00         0,29 Hs.100000         S100 calcium-binding protein of RGS16           4901 TACCTGCAGAA         2,00         0,15 Hs.99519         ESTs           4902 CCAAATGATGA         1,00         0,15 Hs.99519         ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.99519         ESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.89434         drebrin 1           4905 CAACTATCCGT         1,00         0,15 Hs.87595         translocase of inner mitochondrial membrane 22 (yeas           4906 GCTTCATCAGA         1,00         0,15 Hs.75193         COP9 homolog           4908 GCTGGAGAGAG         1,00         0,15 Hs.6396         jumpling translocation breakpoint           4909 GGGCAGAGAGA         1,00         0,15 Hs.6278<	4895	TGAACAGTAAT	2,00	0,29	Hs.80120	UDP-N-acetyl-alpha-D-
4897 AACAACTGGCT         2,00         0,29 Hs.75243         bromodomain-containing 2           4898 CAGGTCAAGAA         2,00         0,29 Hs.179565         minichromosome maintenance deficient (S. cerevisiae)           4899 CTGCCTCCGTG         2,00         0,29 Hs.12851         ESTs, Weakly similar to PSS1_HUMAN PHOSPHATIDYLSERIN           4900 ATTACACCACG         2,00         0,29 Hs.107014         membrane interacting protein of RGS16           4901 TACCTGCAGAA         2,00         0,29 Hs.100000         S100 calcium-binding protein A8 (calgranulin A)           4902 CCAAATGATGA         1,00         0,15 Hs.99519         ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.99519         ESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.89434         drebrin 1           4905 CAACTATCCGT         1,00         0,15 Hs.87595         retinoblastoma-binding protein 2           4906 GCTTCATCAGA         1,00         0,15 Hs.76272         retinoblastoma-binding protein 2           4907 ATACACTTTGT         1,00         0,15 Hs.63432         hypothetical protein FLJ10913           4909 GGGCAGAGAGA         1,00         0,15 Hs.6278         DKFZP586B1621 protein           4910 GAAGCAAGAAA         1,00         0,15 Hs.285027         LIM domain kinase 2           4912 TGCAGGTACTG         1,00						galactosamine:polypeptide N-ace
4898 CAGGTCAAGAA         2,00         0,29 Hs.179565 minichromosome maintenance deficient (S. cerevisiae)           4899 CTGCCTCCGTG         2,00         0,29 Hs.12851 ESTs, Weakly similar to PSS1_HUMAN PHOSPHATIDYLSERIN           4900 ATTACACCACG         2,00         0,29 Hs.107014 membrane interacting protein of RGS16           4901 TACCTGCAGAA         2,00         0,29 Hs.100000 S100 calcium-binding protein A8 (calgranulin A)           4902 CCAAATGATGA         1,00         0,15 Hs.99519 ESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.89434 drebrin 1           4905 CAACTATCCGT         1,00         0,15 Hs.87595 translocase of inner mitochondrial membrane 22 (yeas           4906 GCTTCATCAGA         1,00         0,15 Hs.76272 retinoblastoma-binding protein 2           4907 ATACACTTTGT         1,00         0,15 Hs.64322 hypothetical protein FLJ10913 jumping translocation breakpoint           4908 GCTTGGAGTTA         1,00         0,15 Hs.6396 jumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.23260 presenilin 1 (Alzheimer disease 3)           4911 TGCAGGTCACA         1,00         0,15 Hs.278027 LIM domain kinase 2           4913 GGGAAATTGG         1,00         0,15 Hs.24125 putative zinc finger protein           4914 GAAATCCCAAT         1,00         0,15 Hs.24062 hypothetical protein           4915 GGAAGAAGAAA         1,00	4896	TCCAGAATCCT	2,00	0,29	Hs.7764	
S. cerevisiae    (S. cerevisiae)   (S. cerevis						
PHOSPHATIDYLSERIN	4898	CAGGTCAAGAA	2,00	0,29	Hs.179565	
4901 TACCTGCAGAA         2,00         0,29 Hs.100000 S100 calcium-binding protein A8 (calgranulin A)           4902 CCAAATGATGA         1,00         0,15 Hs.99519 ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.89434 IESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.89434 Iranslocase of inner mitochondrial membrane 22 (yeas           4905 CAACTATCCGT         1,00         0,15 Hs.76272 Iranslocase of inner mitochondrial membrane 22 (yeas           4906 GCTTCATCAGA         1,00         0,15 Hs.75193 Icope homolog           4908 GCTTGGAGTTA         1,00         0,15 Hs.64322 Iranslocation breakpoint           4909 GGGCAGAGAGA         1,00         0,15 Hs.6396 Illumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.3260 Illumping translocation breakpoint           4911 TGCAGGTACTG         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4913 GGGGAAATTGG         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4914 GAAATCCCAAT         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4915 GGGGAAGAGAAA         1,00         0,15 Hs.224062 Injunction protein FLJ10563           4916 GAGGAGGTGGA         1,00         0,15 Hs.224062 Injunction protein pr	4899	CTGCCTCCGTG	2,00	0,29	Hs.12851	
4901 TACCTGCAGAA         2,00         0,29 Hs.100000 S100 calcium-binding protein A8 (calgranulin A)           4902 CCAAATGATGA         1,00         0,15 Hs.99519 ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.89434 IESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.89434 Iranslocase of inner mitochondrial membrane 22 (yeas           4905 CAACTATCCGT         1,00         0,15 Hs.76272 Iranslocase of inner mitochondrial membrane 22 (yeas           4906 GCTTCATCAGA         1,00         0,15 Hs.75193 Icope homolog           4908 GCTTGGAGTTA         1,00         0,15 Hs.64322 Iranslocation breakpoint           4909 GGGCAGAGAGA         1,00         0,15 Hs.6396 Illumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.3260 Illumping translocation breakpoint           4911 TGCAGGTACTG         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4913 GGGGAAATTGG         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4914 GAAATCCCAAT         1,00         0,15 Hs.278027 Illumping translocation breakpoint           4915 GGGGAAGAGAAA         1,00         0,15 Hs.224062 Injunction protein FLJ10563           4916 GAGGAGGTGGA         1,00         0,15 Hs.224062 Injunction protein pr	4900	ATTACACCACG	2,00	0,29	Hs.107014	membrane interacting protein of RGS16
(calgranulin A)	4901	TACCTGCAGAA				
4902 CCAAATGATGA         1,00         0,15 Hs.99519         ESTs           4903 TCACGCGCTCC         1,00         0,15 Hs.93231         ESTs           4904 CTGGGATGCAG         1,00         0,15 Hs.89434         drebrin 1           4905 CAACTATCCGT         1,00         0,15 Hs.87595         translocase of inner mitochondrial membrane 22 (yeas           4906 GCTTCATCAGA         1,00         0,15 Hs.76272         retinoblastoma-binding protein 2           4907 ATACACTTTGT         1,00         0,15 Hs.75193         COP9 homolog           4908 GCTTGGAGTTA         1,00         0,15 Hs.64322         hypothetical protein FLJ10913           4909 GGGGCAGAGAG         1,00         0,15 Hs.6396         jumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.6278         DKFZP586B1621 protein           4911 TTGAAGTCAAA         1,00         0,15 Hs.278027         LIM domain kinase 2           4912 TGCAGGTACTG         1,00         0,15 Hs.265960         hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.265960         hypothetical protein FLJ10563           4915 GGAGGAGGAGAA         1,00         0,15 Hs.224062         hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.20625         sin3-associated polypeptide,				·		
4904         CTGGGATGCAG         1,00         0,15         Hs.89434         drebrin 1           4905         CAACTATCCGT         1,00         0,15         Hs.87595         translocase of inner mitochondrial membrane 22 (yeas           4906         GCTTCATCAGA         1,00         0,15         Hs.76272         retinoblastoma-binding protein 2           4907         ATACACTTTGT         1,00         0,15         Hs.75193         COP9 homolog           4908         GCTTGGAGTTA         1,00         0,15         Hs.64322         hypothetical protein FLJ10913           4909         GGGGCAGAGAG         1,00         0,15         Hs.6396         jumping translocation breakpoint           4910         GAAGCAAGAAA         1,00         0,15         Hs.6278         DKFZP586B1621 protein           4911         TTGAAGTCAAA         1,00         0,15         Hs.278027         LIM domain kinase 2           4912         TGCAGGTACTG         1,00         0,15         Hs.265960         hypothetical protein FLJ10563           4914         GAAATCCCAAT         1,00         0,15         Hs.24125         putative zinc finger protein           4915         GGAGGAGAGAA         1,00         0,15         Hs.24062         hypothetical protein           49	4902	CCAAATGATGA	1,00	0,15	Hs.99519	ESTs
4904         CTGGGATGCAG         1,00         0,15         Hs.89434         drebrin 1           4905         CAACTATCCGT         1,00         0,15         Hs.87595         translocase of inner mitochondrial membrane 22 (yeas           4906         GCTTCATCAGA         1,00         0,15         Hs.76272         retinoblastoma-binding protein 2           4907         ATACACTTTGT         1,00         0,15         Hs.75193         COP9 homolog           4908         GCTTGGAGTTA         1,00         0,15         Hs.64322         hypothetical protein FLJ10913           4909         GGGGCAGAGAG         1,00         0,15         Hs.6396         jumping translocation breakpoint           4910         GAAGCAAGAAA         1,00         0,15         Hs.6278         DKFZP586B1621 protein           4911         TTGAAGTCAAA         1,00         0,15         Hs.26278         DKFZP586B1621 protein           4912         TGCAGGTACTG         1,00         0,15         Hs.278027         LlM domain kinase 2           4913         GGGGAAATTGG         1,00         0,15         Hs.265960         hypothetical protein FLJ10563           4914         GAAATCCCAAT         1,00         0,15         Hs.241052         putative zinc finger protein	4903	TCACGCGCTCC	1,00			ESTs
4905         CAACTATCCGT         1,00         0,15         Hs.87595         translocase of inner mitochondrial membrane 22 (yeas           4906         GCTTCATCAGA         1,00         0,15         Hs.76272         retinoblastoma-binding protein 2           4907         ATACACTITGT         1,00         0,15         Hs.75193         COP9 homolog           4908         GCTTGGAGTTA         1,00         0,15         Hs.64322         hypothetical protein FLJ10913           4909         GGGGCAGAGAG         1,00         0,15         Hs.6396         jumping translocation breakpoint           4910         GAAGCAAGAAA         1,00         0,15         Hs.6278         DKFZP586B1621 protein           4911         TTGAAGTCAAA         1,00         0,15         Hs.3260         presenilin 1 (Alzheimer disease 3)           4912         TGCAGGTACTG         1,00         0,15         Hs.278027         LIM domain kinase 2           4913         GGGGAAATTGG         1,00         0,15         Hs.265960         hypothetical protein FLJ10563           4914         GAAATCCCAAT         1,00         0,15         Hs.24125         putative zinc finger protein           4915         GGAGAGAGAAA         1,00         0,15         Hs.224062         hypothetical protein <td>4904</td> <td>CTGGGATGCAG</td> <td>1,00</td> <td>0,15</td> <td>Hs.89434</td> <td>drebrin 1</td>	4904	CTGGGATGCAG	1,00	0,15	Hs.89434	drebrin 1
4906         GCTTCATCAGA         1,00         0,15         Hs.76272         retinoblastoma-binding protein 2           4907         ATACACTTTGT         1,00         0,15         Hs.75193         COP9 homolog           4908         GCTTGGAGTTA         1,00         0,15         Hs.64322         hypothetical protein FLJ10913           4909         GGGCAGAGAG         1,00         0,15         Hs.6396         jumping translocation breakpoint           4910         GAAGCAAGAAA         1,00         0,15         Hs.6278         DKFZP586B1621 protein           4911         TTGAAGTCAAA         1,00         0,15         Hs.3260         presenilin 1 (Alzheimer disease 3)           4912         TGCAGGTACTG         1,00         0,15         Hs.278027         LIM domain kinase 2           4913         GGGGAAATTGG         1,00         0,15         Hs.265960         hypothetical protein FLJ10563           4914         GAAATCCCAAT         1,00         0,15         Hs.24125         putative zinc finger protein           4915         GGAGAGAGAAA         1,00         0,15         Hs.24062         hypothetical protein           4917         TAGAAATGTTC         1,00         0,15         Hs.29085         sin3-associated polypeptide, 30kD <t< td=""><td>4905</td><td>CAACTATCCGT</td><td>1,00</td><td></td><td></td><td>translocase of inner mitochondrial</td></t<>	4905	CAACTATCCGT	1,00			translocase of inner mitochondrial
4907 ATACACTTTGT         1,00         0,15 Hs.75193         COP9 homolog           4908 GCTTGGAGTTA         1,00         0,15 Hs.64322         hypothetical protein FLJ10913           4909 GGGCAGAGAG         1,00         0,15 Hs.6396         jumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.6278         DKFZP586B1621 protein           4911 TTGAAGTCAAA         1,00         0,15 Hs.3260         presenilin 1 (Alzheimer disease 3)           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 LIM domain kinase 2           4913 GGGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.1230						membrane 22 (yeas
4908 GCTTGGAGTTA         1,00         0,15 Hs.64322         hypothetical protein FLJ10913           4909 GGGGCAGAGAG         1,00         0,15 Hs.6396         jumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.6278         DKFZP586B1621 protein           4911 TTGAAGTCAAA         1,00         0,15 Hs.3260         presenilin 1 (Alzheimer disease 3)           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 LIM domain kinase 2           4913 GGGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein					Hs.76272	retinoblastoma-binding protein 2
4909 GGGCAGAGAG         1,00         0,15 Hs.6396         jumping translocation breakpoint           4910 GAAGCAAGAAA         1,00         0,15 Hs.6278         DKFZP586B1621 protein           4911 TTGAAGTCAAA         1,00         0,15 Hs.3260         presenilin 1 (Alzheimer disease 3)           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 LIM domain kinase 2           4913 GGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.29985 sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGAGGGAG         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGAGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein	4907	ATACACTTTGT				COP9 homolog
4910 GAAGCAAGAAA         1,00         0,15 Hs.6278         DKFZP586B1621 protein           4911 TTGAAGTCAAA         1,00         0,15 Hs.3260         presenilin 1 (Alzheimer disease 3)           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 LlM domain kinase 2           4913 GGGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985 sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein	4908	GCTTGGAGTTA				hypothetical protein FLJ10913
4911 TTGAAGTCAAA         1,00         0,15 Hs.3260         presenilin 1 (Alzheimer disease 3)           4912 TGCAGGTACTG         1,00         0,15 Hs.278027 LIM domain kinase 2           4913 GGGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985 sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein						jumping translocation breakpoint
4912 TGCAGGTACTG         1,00         0,15 Hs.278027 LIM domain kinase 2           4913 GGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985 sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein						
4913 GGGGAAATTGG         1,00         0,15 Hs.265960 hypothetical protein FLJ10563           4914 GAAATCCCAAT         1,00         0,15 Hs.24125 putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985 sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein			1,00			
4914 GAAATCCCAAT         1,00         0,15 Hs.24125         putative zinc finger protein           4915 GGAAGAAGAAA         1,00         0,15 Hs.240062         hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466         ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985         sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525         ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305         DKFZP566B183 protein						
4915 GGAAGAAGAAA         1,00         0,15 Hs.240062 hypothetical protein           4916 GAGGAGGTGGA         1,00         0,15 Hs.22466 ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985 sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525 ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein	4913	GGGGAAATTGG	1,00			
4916 GAGGAGGTGGA         1,00         0,15 Hs.22466         ubiquitin specific protease 19           4917 TAGAAATGTTC         1,00         0,15 Hs.20985         sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525         ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305         DKFZP566B183 protein			1,00	0,15	Hs.24125	putative zinc finger protein
4917 TAGAAATGTTC         1,00         0,15 Hs.20985         sin3-associated polypeptide, 30kD           4918 ATGACTGCTGT         1,00         0,15 Hs.19525         ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein			1,00			
4918 ATGACTGCTGT         1,00         0,15 Hs.19525         ESTs           4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein			1,00			
4919 GCATTCGCAGT         1,00         0,15 Hs.183842 ubiquitin B           4920 TTACACAGACT         1,00         0,15 Hs.182447 heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921 CTGGGAGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein	4917	TAGAAATGTTC	1,00	0,15	Hs.20985	sin3-associated polypeptide, 30kD
4920         TTACACAGACT         1,00         0,15         Hs.182447         heterogeneous nuclear ribonucleoprotein C (C1/C2)           4921         CTGGGAGGGAG         1,00         0,15         Hs.154304         Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922         TTGTGTTCTTT         1,00         0,15         Hs.12305         DKFZP566B183 protein			1,00			
C (C1/C2)  4921 CTGGGAGGGAG 1,00 0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19  4922 TTGTGTTCTTT 1,00 0,15 Hs.12305 DKFZP566B183 protein			1,00	0,15	Hs.183842	ubiquitin B
4921 CTGGGAGGAG         1,00         0,15 Hs.154304 Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19           4922 TTGTGTTCTTT         1,00         0,15 Hs.12305 DKFZP566B183 protein	4920	TTACACAGACT	1,00	0,15	Hs.182447	
4922 TTGTGTTCTTT 1,00 0,15 Hs.12305 DKFZP566B183 protein	4921	CTGGGAGGGAG	1,00	0,15	Hs.154304	Homo sapiens chromosome 19, BAC
	4922	TIGIGTICTIT	1,00	0,15	Hs.12305	
			1,00			

	1921	AGCAGCCTTTT	1,00	0.15	He 10496	craniofacial development protein 1
4926 CTTCTACTAAT         6,00         0,55 Hs.109857 Homo sapiens mRNA; cDNA DKFZP434H0820 (from clone DK           4927 CGCGTGCACAC         4,00         0,45 Hs.104640 HIV-1 inducer of short transcripts binding protein           4928 GTAGCAGGGCT         3,00         0,38 Hs.32797         ESTs           4929 TGCAGAACGGC         3,00         0,38 Hs.20776         CGI-91 protein           4931 TGGCAATGAA         7,00         0,59 Hs.8170         hypothetical protein           4931 TGGGCAGCAG         4,00         0,45 Hs.13743         ESTs, Weakly similar to A28996 proline-rich protein           4932 CCAGGGTGCGT         4,00         0,45 Hs.149846 integrin, beta 5         4933 TTTCTGCTCCT         4,00         4,54 Hs.178869         transcription elongation factor A (SII), 1           4933 TTCTGCACTCT         4,00         0,29 Hs.78869         transcription elongation factor A (SII), 1           4936 GGTGTATAATT         2,00         0,29 Hs.20529         ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.20013         DKFZP564G2022 protein           4938 CCACTTITTAA         2,00         0,29 Hs.11649         DKFZP564G2022 protein           4938 CACTCGCTCTA         2,00         0,29 Hs.110418 ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179419         Hs.17919 Iprotein </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
By   By   By   By   By   By   By   By						
4927 CGCGTGCACAC         4,00         0,45 Hs.104640 HIV-1 inducer of short transcripts binding protein           4928 GTAGCAGGGCT         3,00         0,38 Hs.32797 ESTs           4929 TGCAGAACGGC         3,00         0,38 Hs.20776 CGI-91 protein           4930 TCTGCAATGAA         7,00         0,59 Hs.8170 Nypothetical protein           4931 TGAGTGGACAG         4,00         0,45 Hs.3743 Nypothetical protein           4931 TGAGTGGACAG         4,00         0,45 Hs.13743 Nypothetical protein           4932 CCAGGCTGCGT         4,00         0,45 Hs.14846 Integrin, beta 5           4933 TTTCTGCTCCT         4,00         0,45 Hs.18424 libosomal protein L41           4934 CTTTGCACTCT         2,00         0,29 Hs.78869 transcription elongation factor A (SII), 1           4936 GGTGAATATT         2,00         0,29 Hs.20529 ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.2013 DKFZP564Q2082 protein           4938 CACTTTTTAA         2,00         0,29 Hs.11615 map kinase phosphatase-like protein MK-STYX           4940 GTTTCCGGTT         2,00         0,29 Hs.111449 CGI-94 protein           4941 AGTATGCCACT         2,00         0,29 Hs.119419 CGI-94 protein           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type 1 protein FLJ111342           4943 GCCGTGAGGACA         7	4920	CITCIACIAAI	0,00	0,55	ns. 109657	
Protein   Prot	4007	CCCCTCCACAC	4.00	0.45	Ho 104640	
4928 GTACCAGGCT         3,00         0,38 Hs. 20776         ESTS           4929 TGCAGAACGC         3,00         0,38 Hs. 20776         CGI-91 protein           4930 TCTGCAATGAA         7,00         0,59 Hs. 8170         hypothetical protein           4931 TGAGTGGACAG         4,00         0,45 Hs. 3743         ESTS, Weakly similar to A28996 proline-rich protein           4932 CCAGGCTGCGT         4,00         0,45 Hs. 149846 integrin, beta 5           4933 TTTCTGCACTCT         4,00         0,45 Hs. 108124 inbosomal protein L41           4934 CTTTGCACTCT         2,00         0,29 Hs. 74170         metallothionein 1E (functional)           4935 GCTGTATAATT         2,00         0,29 Hs. 20529         ESTS, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs. 20013         DKFZP564C2082 protein           4937 TACAGTTCCCT         2,00         0,29 Hs. 16492         DKFZP564C2082 protein           4938 CCACTTTITTAA         2,00         0,29 Hs. 11615         map kinase phosphatase-like protein           4940 GTTTCCGGTT         2,00         0,29 Hs. 110418         ESTs           4941 AGTATGCCACT         2,00         0,29 Hs. 179516 integral type I protein           4943 GCCGTGAGCAG         7,00         0,59 Hs. 779516 integral type I protein           494	4921	CGCGTGCACAC	4,00	0,45		
1929   TGCAGAACGGC   3,00   0,38   Hs. 20776   CGI-91 protein   1930   TCTGCAATGAA   7,00   0,59   Hs. 3170   hypothetical protein   1931   TGAGTGGACAG   4,00   0,45   Hs. 3743   ESTs, Weakly similar to A28996 proline-rich protein   1932   TCTGCTCCT   4,00   0,45   Hs. 149846   integrin, beta 5   1933   TTTCTGCTCCT   4,00   0,29   Hs. 78869   transcription elongation factor A (SII), 1   1934   CTTTGCACTCT   2,00   0,29   Hs. 78869   transcription elongation factor A (SII), 1   1935   GCTGTATAATT   2,00   0,29   Hs. 784170   metallothionein 1E (functional)   1936   GGTTGAGTGG   2,00   0,29   Hs. 20013   DKFZP56402082 protein   1938   CCACTTTITAA   2,00   0,29   Hs. 10418   ESTs   1939   GATCCGCTCTA   2,00   0,29   Hs. 11615   map kinase phosphatase-like protein   1944   AGTATGCCACT   2,00   0,29   Hs. 11449   CGI-94 protein   1944   AGTATGCCACT   2,00   0,29   Hs. 11418   ESTs   1942   CAAGCAGGACA   7,00   0,59   Hs. 179516   integral type I protein   1943   GCCGTGAGCAG   3,00   0,38   Hs. 266514   hypothetical protein   FLJ11342   1944   AGATGAGACC   10,00   0,70   Hs. 183698   ribosomal protein   L29   1949   GACTCAGGAAA   15,00   0,39   Hs. 77516   high density lipoprotein binding protein   1949   GACTCAGCACC   2,00   0,29   Hs. 30418   Hs. 3	4020	CTACCACCCCT	2.00	0.20	Un 22707	
1930   TCTGCAATGAA   7,00   0,59   Hs. 8170   hypothetical protein   1931   TGAGTGGACAG   4,00   0,45   Hs. 3743   ESTs, Weakly similar to A28996 proline-rich protein   1932   CCAGGCTGCGT   4,00   0,45   Hs. 149846   integrin, beta 5   1933   TTTCTGCTCCT   4,00   0,45   Hs. 18124   fibosomal protein L41   1934   CTTTGCACTCT   2,00   0,29   Hs. 78899   transcription elongation factor A (SII), 1   1935   GCTGTATAATT   2,00   0,29   Hs. 20529   ESTs, Weakly similar to ubiquitous TPR motif, Y isof   1937   TACAGTTCCCT   2,00   0,29   Hs. 20529   ESTs, Weakly similar to ubiquitous TPR motif, Y isof   1938   CCACTTTTAA   2,00   0,29   Hs. 16492   DKFZP564G2022 protein   1938   GATCCGCTCTA   2,00   0,29   Hs. 16492   DKFZP564G2022 protein   1938   GATCCGCTCTA   2,00   0,29   Hs. 11615   Ms. STYX   1949   Ms. STYX						
4931 TGAGTGGACAG         4,00         0,45 Hs.3743         ESTs, Weakly similar to A28996 proline-rich protein           4932 CCAGGCTGCGT         4,00         0,45 Hs.149846 lintegrin, beta 5           4933 TTTCTGCTCT         4,00         0,45 Hs.108124 ribosomal protein L41           4935 GCTGTATAATT         2,00         0,29 Hs.78869 transcription elongation factor A (SII), 1           4936 GGTGAGTGTG         2,00         0,29 Hs.20529 ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.20013 DKFZP56402082 protein           4938 CCACTTITTAA         2,00         0,29 Hs.16492 DKFZP564G2022 protein           4938 CACTTTTTAA         2,00         0,29 Hs.111449 CGI-94 protein           4940 GTTTTCCGGTT         2,00         0,29 Hs.110418 ESTs           4941 AGTATGCCACT         2,00         0,29 Hs.110418 ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACG         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigillin)           4946 GGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)						
4932 CCAGGCTGCGT         4,00         0,45 Hs.149846 integrin, beta 5           4933 TTTCTGCTCCT         4,00         0,45 Hs.108124 ribosomal protein L41           4934 CTTTGCACTCT         2,00         0,29 Hs.78869         transcription elongation factor A (SII), 1           4935 GCTGTATAATT         2,00         0,29 Hs.20529         ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.20013         DKFZP564C2082 protein           4938 CCACTTTTAA         2,00         0,29 Hs.16492         DKFZP564G2022 protein           4939 GATCCGCTCTA         2,00         0,29 Hs.11615         map kinase phosphatase-like protein MK-STYX           4940 GTTTCCGGTT         2,00         0,29 Hs.111449 CGI-94 protein           4941 AGTATGCCACT         2,00         0,29 Hs.110418 ESTs           4942 CAAGCAGGACA         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACC         10,00         0,70 Hs.183698 ribosomal protein L29           4945 ACCTCAGCACA         3,00         0,39 Hs.77171           4948 GGACTTTCCTT         3,00         0,39 Hs.75789         N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.63525	4931	IGAGIGGACAG	4,00	0,45	HS.3/43	
4933 TTTCTGCTCCT         4,00         0,45 Hs.108124 ribosomal protein L41           4934 CTTTGCACTCT         2,00         0,29 Hs.78869 transcription elongation factor A (SII), 1           4935 GCTGTATAATT         2,00         0,29 Hs.74170 metallothionein 1E (functional)           4936 GGTGAGTGTG         2,00         0,29 Hs.20529 ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.20013 DKFZP564G2028 protein           4938 CCACTTTTAA         2,00         0,29 Hs.16492 DKFZP564G2022 protein           4939 GATCCGCTCTA         2,00         0,29 Hs.11615 map kinase phosphatase-like protein MK-STYX           4940 GTTTCCGGTT         2,00         0,29 Hs.110418 ESTs           4941 AGTATGCCACT         2,00         0,29 Hs.179516 integral type I protein           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 BAGTGAGACCC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTAGCACA         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S.	4022	COACCOTOCCT	4.00	0.45	U- 140046	
4934 CTTTGCACTCT         2,00         0,29 Hs.78869         transcription elongation factor A (SII), 1           4935 GCTGTATAATT         2,00         0,29 Hs.20529         ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.20013         DKFZP564O2082 protein           4938 CCACTTITTAA         2,00         0,29 Hs.16492         DKFZP564G2022 protein           4939 GATCCGCTCTA         2,00         0,29 Hs.11615         MKFZP564G2022 protein           4940 GTTTCCGGTT         2,00         0,29 Hs.110418         ESTs           4941 AGTATGCCACT         2,00         0,29 Hs.110418         ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACCC         10,00         0,70 Hs.125359         Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516         high density lipoprotein binding protein (vigilin)           4946 GGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.75789         N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.3566         ART-4 protein						
4935 GCTGTATAATT         2,00         0,29 Hs.74170         metallothionein 1E (functional)           4936 GGTTGAGTGTG         2,00         0,29 Hs.20529         ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937 TACAGTTCCCT         2,00         0,29 Hs.20013         DKFZP564C2082 protein           4938 CCACTTTTAA         2,00         0,29 Hs.16492         DKFZP564G2022 protein           4940 GTTTCCGGTT         2,00         0,29 Hs.11615         map kinase phosphatase-like protein MK-STYX           4940 GTTTCCGGTT         2,00         0,29 Hs.110418         ESTs           4941 AGTATGCCACT         2,00         0,29 Hs.110418         ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hybothetical protein FLJ11342           4944 GAGTGAGACC         10,00         0,70 Hs.183698 ribosomal protein L29           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigillin)           4946 GGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACACACCC         3,00         0,39 Hs.75789         N-myc downstream regulated major histocompatibility complex, class lil. DP alpha           4948 GGACTTTCCTT         3,00         0,29 Hs.63525         N-my						
4936         GGTTGAGTGTG         2,00         0,29 Hs.20529         ESTs, Weakly similar to ubiquitous TPR motif, Y isof           4937         TACAGTTCCCT         2,00         0,29 Hs.20013         DKFZP564O2082 protein           4938         CCACTTTTAA         2,00         0,29 Hs.16492         DKFZP564G2022 protein           4939         GATCCGCTCTA         2,00         0,29 Hs.11615         map kinase phosphatase-like protein MK-STYX           4940         GTTTCCGGTT         2,00         0,29 Hs.110418         ESTs           4941         AGTATGCCACT         2,00         0,29 Hs.110418         ESTs           4942         CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943         GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944         GAGTGAGACC         10,00         0,70 Hs.183698 ribosomal protein L29 uninichromosome maintenance deficient (vigillin)           4946         GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947         GACTCGCCCAC         3,00         0,39 Hs.75789         N-myc downstream regulated           4949         TGAAAACTACT         2,00         0,29 Hs.63525         N-Myc downstream regulated           4949         TGAATGATTTAC         2						
Motif, Y isof   Motif, Y iso						
4937 TACAGTTCCCT         2,00         0,29 Hs.20013         DKFZP564O2082 protein           4938 CCACTTTTAA         2,00         0,29 Hs.16492         DKFZP564G2022 protein           4939 GATCCGCTCTA         2,00         0,29 Hs.11615         map kinase phosphatase-like protein           4940 GTTTCCGGTT         2,00         0,29 Hs.110418         ESTs           4941 AGTATGCCACT         2,00         0,29 Hs.110418         ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACCC         10,00         0,70 Hs.125359         Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)         (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29         N-myc downstream regulated           4948 GGACTTTCCTT         3,00         0,39 Hs.75789         N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.34578         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951 TGGGTTTTAAA         2,00         0,29 Hs.3566         ART-4 protein           4952 CATTTAGATTT         2,00         0,29 Hs.3566	4936	GGTGAGTGTG	2,00	0,29	Hs.20529	
4938 CCACTTTTAA         2,00         0,29 Hs.16492         DKFZP564G2022 protein           4939 GATCCGCTCTA         2,00         0,29 Hs.11615         map kinase phosphatase-like protein           4940 GTTTCCGGTT         2,00         0,29 Hs.111449         CGI-94 protein           4941 AGTATGCCACT         2,00         0,29 Hs.110418         ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171         minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789         N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.74578         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951 TGGGTTTTAAA         2,00         0,29 Hs.63525         poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.270605         ESTs, Weakly similar t	1007	T1010T00T	0.00	0.00		
4939 GATCCGCTCTA         2,00         0,29 Hs.11615         map kinase phosphatase-like protein MK-STYX           4940 GTTTTCCGGTT         2,00         0,29 Hs.111449 CGI-94 protein           4941 AGTATGCCACT         2,00         0,29 Hs.110418 ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACCC         10,00         0,70 Hs.125359           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789 N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.74578 DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4950 CTATGTGTTAC         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4951 TGGGTTTAAA         2,00         0,29 Hs.3566 ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4954 TGAATGATTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence						
MK-STYX						
4940 GTTTTCCGGTT         2,00         0,29 Hs.111449 CGI-94 protein           4941 AGTATGCCACT         2,00         0,29 Hs.110418 ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACCC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789 N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914 major histocompatibility complex, class III, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4951 TGGGTTTAAA         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.3560 ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4954 TGAATGATTTT         2,00         0,29 Hs.192707 glioblastoma amplified sequence      <	4939	GATCCGCTCTA	2,00	0,29		
4941 AGTATGCCACT         2,00         0,29 Hs.110418 ESTs           4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACCC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789 nyc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914 major histocompatibility complex, class III, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4951 TGGGTTTAAA         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.270605 ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.270605 ESTs           4955 GCCAGAAGGGG         1,00         0,15 Hs.8984 Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072 ESTs						
4942 CAAGCAGGACA         7,00         0,59 Hs.179516 integral type I protein           4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789 N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914 major histocompatibility complex, class II, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4951 TGGGTTTAAA         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.270605 ESTs, Weakly similar to alternatively spliced produc           4953 CCTGGAATCCC         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.8984 Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072 ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.7			2,00			
4943 GCCGTGAGCAG         3,00         0,38 Hs.266514 hypothetical protein FLJ11342           4944 GAGTGAGACC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789 N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914 major histocompatibility complex, class II, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4951 TGGGTTTAAA         2,00         0,29 Hs.3566 ART-4 protein           4952 CATTTAGATTT         2,00         0,29 Hs.270605 ESTs, Weakly similar to alternatively spliced produc           4953 CCTGGAATCCC         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825 ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.89072 ESTs           4957 CTCTTATTTCA         1,00         0,15 Hs.89072 ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297 G protein-coupled receptor kinase 6						
4944 GAGTGAGACCC         10,00         0,70 Hs.125359 Thy-1 cell surface antigen           4945 ACCTCAGGAAA         15,00         0,84 Hs.177516 high density lipoprotein binding protein (vigilin)           4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTCCTT         3,00         0,39 Hs.75789 N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914 major histocompatibility complex, class II, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.74578 DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951 TGGGTTTAAA         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.3566 ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.270605 ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.8984 Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072 ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.75232 SEC14 (S. cerevisiae)-like 1           4959 ACCGCACAAAT         1,00 <t< td=""><td></td><td></td><td></td><td></td><td></td><td></td></t<>						
4945         ACCTCAGGAAA         15,00         0,84         Hs.177516         high density lipoprotein binding protein (vigilin)           4946         GGGCTGGGGTC         100,00         2,07         Hs.183698         ribosomal protein L29           4947         GACTCGCCCAC         3,00         0,39         Hs.77171         minichromosome maintenance deficient (S. cerevisiae)           4948         GGACTTTCCTT         3,00         0,39         Hs.75789         N-myc downstream regulated           4949         TGAAAACTACT         2,00         0,29         Hs.914         major histocompatibility complex, class II, DP alpha           4950         CTATGTGTTAC         2,00         0,29         Hs.63525         poly(rC)-binding protein 2           4951         TGGGTTTTAAA         2,00         0,29         Hs.3566         ART-4 protein           4952         CATTTAGATTT         2,00         0,29         Hs.270605         ESTs, Weakly similar to alternatively spliced produc           4954         TGAATGATTTT         2,00         0,29         Hs.152707         glioblastoma amplified sequence           4955         GCCAGAAGGGG         1,00         0,15         Hs.8984         Homo sapiens chromosome 14 BAC           4957         CTCTTATTTCA         1,00         0,15				0,38	Hs.266514	hypothetical protein FLJ11342
(vigilin)				0,70	Hs.125359	Thy-1 cell surface antigen
4946 GGGCTGGGGTC         100,00         2,07 Hs.183698 ribosomal protein L29           4947 GACTCGCCAC         3,00         0,39 Hs.77171 minichromosome maintenance deficient (S. cerevisiae)           4948 GGACTTTCCTT         3,00         0,39 Hs.75789 N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914 major histocompatibility complex, class II, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.74578 DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA)           4951 TGGGTTTTAAA         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.270605 ESTs, Weakly similar to alternatively spliced produc           4953 CCTGGAATCCC         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825 ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984 Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.76297 G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232 SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838 hypothetical protein PRO0971	4945	ACCTCAGGAAA	15,00	0,84	Hs.177516	
4947         GACTCGCCCAC         3,00         0,39         Hs.77171         minichromosome maintenance deficient (S. cerevisiae)           4948         GGACTTTCCTT         3,00         0,39         Hs.75789         N-myc downstream regulated           4949         TGAAAACTACT         2,00         0,29         Hs.914         major histocompatibility complex, class II, DP alpha           4950         CTATGTGTTAC         2,00         0,29         Hs.74578         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951         TGGGTTTTAAA         2,00         0,29         Hs.63525         poly(rC)-binding protein 2           4952         CATTTAGATTT         2,00         0,29         Hs.3566         ART-4 protein           4953         CCTGGAATCCC         2,00         0,29         Hs.152707         glioblastoma amplified sequence           4954         TGAATGATTTT         2,00         0,15         Hs.90825         ESTs           4955         GCCAGAAGGGG         1,00         0,15         Hs.89072         ESTs           4957         CTCTTATTCA         1,00         0,15         Hs.76297         G protein-coupled receptor kinase 6           4959         ACCGCACAAAT         1,00         0,15         Hs.75232         SEC14 (S. cerevisiae)-like 1 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>						
(S. cerevisiae)						
4948 GGACTTTCCTT         3,00         0,39 Hs.75789         N-myc downstream regulated           4949 TGAAAACTACT         2,00         0,29 Hs.914         major histocompatibility complex, class III, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.74578         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951 TGGGTTTAAA         2,00         0,29 Hs.63525         poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.3566         ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.270605         ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971	4947	GACTCGCCCAC	3,00	0,39	Hs.77171	
4949 TGAAAACTACT         2,00         0,29 Hs.914         major histocompatibility complex, class II, DP alpha           4950 CTATGTGTTAC         2,00         0,29 Hs.74578         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951 TGGGTTTTAAA         2,00         0,29 Hs.63525         poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.3566         ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.270605         ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707         glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971					( )	
II, DP alpha						
4950 CTATGTGTTAC         2,00         0,29 Hs.74578         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 9 (RNA           4951 TGGGTTTAAA         2,00         0,29 Hs.63525         poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.3566         ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.270605         ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971	4949	TGAAAACTACT	2,00	0,29		
polypeptide 9 (RNA   4951 TGGGTTTTAAA   2,00   0,29 Hs.63525   poly(rC)-binding protein 2   4952 CATTTAGATTT   2,00   0,29 Hs.3566   ART-4 protein   4953 CCTGGAATCCC   2,00   0,29 Hs.270605   ESTs, Weakly similar to alternatively spliced produc   4954 TGAATGATTT   2,00   0,29 Hs.152707   glioblastoma amplified sequence   4955 GCCAGAAGGGG   1,00   0,15 Hs.90825   ESTs   4956 AAGAAATGCAA   1,00   0,15 Hs.8984   Homo sapiens chromosome 14 BAC   98L12   4957 CTCTTATTTCA   1,00   0,15 Hs.89072   ESTs   4958 TGTGTGTTAGC   1,00   0,15 Hs.76297   G protein-coupled receptor kinase 6   4959 ACCGCACAAAT   1,00   0,15 Hs.75232   SEC14 (S. cerevisiae)-like 1   4960 TGGCCCTTTCA   1,00   0,15 Hs.70838   hypothetical protein PRO0971						
4951 TGGGTTTTAAA         2,00         0,29 Hs.63525 poly(rC)-binding protein 2           4952 CATTTAGATTT         2,00         0,29 Hs.3566 ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.270605 ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825 ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984 Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072 ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297 G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232 SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838 hypothetical protein PRO0971	4950	CTATGTGTTAC	2,00	0,29	Hs.74578	
4952 CATTTAGATTT         2,00         0,29 Hs.3566         ART-4 protein           4953 CCTGGAATCCC         2,00         0,29 Hs.270605         ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971						polypeptide 9 (RNA
4953 CCTGGAATCCC         2,00         0,29 Hs.270605         ESTs, Weakly similar to alternatively spliced produc           4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971						
Spliced produc						
4954 TGAATGATTTT         2,00         0,29 Hs.152707 glioblastoma amplified sequence           4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971	4953	CCTGGAATCCC	2,00	0,29	Hs.270605	, , ,
4955 GCCAGAAGGGG         1,00         0,15 Hs.90825         ESTs           4956 AAGAAATGCAA         1,00         0,15 Hs.8984         Homo sapiens chromosome 14 BAC 98L12           4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971						<del></del>
4956       AAGAAATGCAA       1,00       0,15       Hs.8984       Homo sapiens chromosome 14 BAC 98L12         4957       CTCTTATTTCA       1,00       0,15       Hs.89072       ESTs         4958       TGTGTGTTAGC       1,00       0,15       Hs.76297       G protein-coupled receptor kinase 6         4959       ACCGCACAAAT       1,00       0,15       Hs.75232       SEC14 (S. cerevisiae)-like 1         4960       TGGCCCTTTCA       1,00       0,15       Hs.70838       hypothetical protein PRO0971			2,00			<del>                                     </del>
98L12     98L12       4957   CTCTTATTTCA   1,00   0,15   Hs.89072   ESTs     4958   TGTGTGTTAGC   1,00   0,15   Hs.76297   G protein-coupled receptor kinase 6     4959   ACCGCACAAAT   1,00   0,15   Hs.75232   SEC14 (S. cerevisiae)-like 1   4960   TGGCCCTTTCA   1,00   0,15   Hs.70838   hypothetical protein PRO0971     4957   4958   4959   495			1,00			ESTs
4957 CTCTTATTTCA         1,00         0,15 Hs.89072         ESTs           4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971	4956	AAGAAATGCAA	1,00	0,15	Hs.8984	
4958 TGTGTGTTAGC         1,00         0,15 Hs.76297         G protein-coupled receptor kinase 6           4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971						1
4959 ACCGCACAAAT         1,00         0,15 Hs.75232         SEC14 (S. cerevisiae)-like 1           4960 TGGCCCTTTCA         1,00         0,15 Hs.70838         hypothetical protein PRO0971	4957	CTCTTATTTCA	1,00			ESTs
4960 TGGCCCTTTCA 1,00 0,15 Hs.70838 hypothetical protein PRO0971			1,00	0,15	Hs.76297	G protein-coupled receptor kinase 6
4960 TGGCCCTTTCA 1,00 0,15 Hs.70838 hypothetical protein PRO0971	4959	ACCGCACAAAT	1,00	0,15	Hs.75232	<del> </del>
				0,15	Hs.70838	

1062	TTCGTATTACA	1,00	0.15	Hs.46743	McKusick-Kaufman syndrome
	TTCTGTGTATA	1,00		Hs.3530	TLS-associated serine-arginine protein
	TGTGACACTGA	1,00		Hs.3273	Ts translation elongation factor,
4904	IGIGACACIGA	1,00	0,13	IU8.2512	mitochondrial
4965	GGCTGAATTTT	1,00	0.15	Hs.31921	KIAA0648 protein
	AAACCTGAGAA	1,00		Hs.29724	ESTs, Weakly similar to CDEP
		.,00	٠, . ٠		[H.sapiens]
4967	TCTTCAGTAGC	1,00	0,15	Hs.2934	ribonucleotide reductase M1 polypeptide
4968	GAGCTCAAGAT	1,00			Homo sapiens mRNA; cDNA
		·	,		DKFZp434A012 (from clone DKF
	TATGGGGTCAC	1,00			hypothetical protein
	ATGATGCCTCT	1,00	0,15		KIAA0802 protein
4971	GTTAAACCCTG	1,00	0,15		ESTs, Weakly similar to KIAA0004
1070					[H.sapiens]
	TAATAAGCACT	1,00			MAX binding protein
	GGAGGGGAGGC	1,00			CGI-63 protein
4974	TTATCCTTCAC	1,00	0,15	Hs.188882	Homo sapiens clone 23872 mRNA
4075				100100	sequence
	AATATTAAGAC	1,00		Hs.168103	prp28, U5 snRNP 100 kd protein
	AAGTGCATTTG	1,00		Hs.167420	
4977	TAATGAAATTG	1,00	0,15	Hs.15020	homolog of mouse quaking QKI (KH
4079	CACCTCATCCC	4 00	0.45	11- 405070	domain RNA binding
	CACCTCATCCC	1,00	0,15	HS. 125078	ornithine decarboxylase antizyme 1
4979	AACAAATTCTT	1,00	0,15		fucosyltransferase 8 (alpha (1,6) fucosyltransferase
4980	GGGCTTGGTAT	1,00	0.15		hypothetical protein FLJ10659
	GGCTGGGGGCC	50,00	1 47	Hs.75721	profilin 1
	TATGTGATTTG	3,00	0.30	He 5216	HSPC028 protein
	ACAACACTACA	3,00	0,30	He 277//	RAB3A, member RAS oncogene family
	ATGGCCATAGA	3,00	0,39	Hs 155206	serine/threonine kinase 25 (Ste20, yeast
1007	711000071171071	3,55	0,00	1 13. 100200	homolog)
4985	AAAGCAGCACA	3,00	0,39	Hs.108802	N-ethylmaleimide-sensitive factor
4986	ACTTGGAGCCG	8,00	0,65	Hs.177656	calmodulin 1 (phosphorylase kinase,
		]		_	delta)
	TGCGCGCCCTG	4,00			hypothetical protein
	AAGATAATGCC	4,00	0,46	Hs.102696	MCT-1 protein
	AGCACGACCCG	2,00		Hs.89434	
	TAGTAGATGCT	2,00			cytochrome c oxidase subunit VIc
	CCTCACTTTCT	2,00		Hs.155560	
	TTAGTTAAGCA	3,00			archain 1
4993	ACATCCTCACC	3,00	0,39	Hs.279554	proteasome (prosome, macropain) 26S
4.5					subunit, non-ATP
4994	TCAGCTGGCCC	3,00	0,39	Hs.256583	interleukin enhancer binding factor 3,
4005	COCTTOO	20.00	4 00	11- 70055	90kD
4995	GCCTTCCAATA	30,00	1,20	Hs.76053	DEAD/H (Asp-Glu-Ala-Asp/His) box
1006	CTCCCCCAAGC	E 00	0.50	Ha 20220F	polypeptide 5 (RNA
	CCCTCCTGGGG	5,00			immunoglobulin heavy contant alpha 1
4331		8,00	0,00	Hs.95867	Homo sapiens EST00098 gene, last
	<u> </u>			L	exon

	GAATCACTGCC	5,00			chromosome 2 open reading frame 1
	TTCTCTCAACT	3,00		Hs.27445	unknown
	TAACCAAACAC	3,00			Homo sapiens partial mRNA for NICE-4 protein, 3' end
5001	TCATTTTCCAA	4,00	0,46	Hs.187958	solute carrier family 6 (neurotransmitter transporte
5002	GAAATTTGAAA	2,00	0,30	Hs.79457	hypothetical protein FLJ20519
5003	TCAAGAAACAT	2,00	0,30	Hs.78580	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 1
5004	CAAATAAAATG	2,00	0,30	Hs.71465	squalene epoxidase
5005	GTTCTCAGGAA	2,00	0,30	Hs.6968	KIAA1460 protein
	GTGTAAATGGA	2,00	0,30	Hs.286131	CGI-101 protein
5007	CCACACCGGTA	2,00			heme oxygenase (decycling) 2
5008	AGGTCTGCCAG	2,00	0,30		aldo-keto reductase family 1, member C2 (dihydrodiol
5009	TAGATGTGATG	1,00	0,15		ESTs, Weakly similar to dJ1108D11.1 [H.sapiens]
5010	TAATAAAACCC	1,00	0,15	Hs.94810	Human DNA sequence from clone RP5- 1170K4 on chromoso
5011	ACCCACCTGCA	1,00	0,15	Hs.9100	ESTs
5012	GAGCCTCATCC	1,00	0,15	Hs.83715	Sjogren syndrome antigen B (autoantigen La)
5013	TGGGAAGAGGG	1,00	0,15		enhancer of filamentation 1 (cas-like docking; Crk-a
5014	CGGCAGGAAAA	1,00	0,15	Hs.78019	Homo sapiens clone 24432 mRNA sequence
5015	ATGTATAATAA	1,00	0,15	Hs.67052	vacuolar protein sorting 26 (yeast homolog)
5016	GGTCCAGCATC	1,00	0,15		KIAA0376 protein
5017	TTTAATCTCAG	1,00		Hs.4104	hypothetical protein
5018	AGATTACTGAT	1,00			ESTs
	TGTTGGGTTCT	1,00		Hs.3628	mitogen-activated protein kinase kinase kinase kinas
5020	CAAGAGAGTAG	1,00	0,15	Hs.30332	glutamine-fructose-6-phosphate transaminase 2
5021	TGACACAGCCA	1,00	0,15		ESTs
5022	CTGAATTCAGT	1,00	0,15		ESTs
5023	AATATTGTCCG	1,00	0,15	Hs.283322	hypothetical protein
	ACTTCACAAAG	1,00	0,15	Hs.27556	ESTs
	GTTTTGTGGGA	1,00			hypothetical protein
5026	TTTCATCCACT	1,00	0,15	Hs.25515	KIAA0362 protein
	GTAAAACAATA	1,00	0,15	Hs.24758	ESTs
5028	TTTCATTAATC	1,00	0,15	Hs.247043	type 1 tumor necrosis factor receptor shedding amino
5029	GGGAAACTCCA	1,00	0,15	Hs.235390	Homo sapiens mRNA; cDNA DKFZp761B101 (from clone DKF
5030	TGCCTGTGAAA	1,00	0.15	Hs.228289	
	ATGCATTGTTT	1,00			tryptophan rich basic protein
	CTGTGATTGTG	1,00			CASP8 and FADD-like apoptosis

	<u> </u>	Γ -		ſ <del></del>	regulator
5033	CCAAAATTCTA	1,00	0.15	He 193700	Homo sapiens mRNA; cDNA
3000		1,00	0, 13	13.133700 	DKFZp586I0324 (from clone DK
5034	GACCTGCACTC	1,00	0.15	He 172278	syntrophin, beta 2 (dystrophin-associated
3034	GACCIGCACIC	1,00	0,13	115.172270	protein A1
5025	CCTCTGGCAGC	1,00	0.15	Uc 166235	ESTs, Moderately similar to
3033	CCTCTGGCAGC	1,00	0, 15		ALU5_HUMAN ALU SUBFAMILY
5026	GCAGAAGCACA	1,00	0.15	Uc 16134	serine/threonine kinase 10
-		1,00			
	GTGATGTGCAC			Hs.141683	
	AAAACCTGAAA	1,00			glycogen synthase kinase 3 alpha
	TATAATAAATA	1,00	·		NADH dehydrogenase (ubiquinone) Fe-S protein 4 (18kD
	GACGACACGAG	55,00			ribosomal protein S28
5041	TGGCCCCCGCC	5,00	0,53	Hs.93649	upstream transcription factor 2, c-fos
					interacting
5042	TAATCCTCAAG	4,00			collagen, type XVIII, alpha 1
5043	GTTCTGGTTTA	11,00	0,77	Hs.241336	Homo sapiens mRNA; cDNA
					DKFZp564G0422 (from clone DK
5044	TAAGTAGCAAA	6,00			integral membrane protein 2B
5045	GCTTACCTTTC	8,00	0,67	Hs.7753	calumenin
5046	CCACTCCTCCA	4,00	0,47	Hs.82890	defender against cell death 1
5047	CAGCGCTTTGA	4,00		Hs.7274	DKFZP434P1750 protein
5048	GTACTCCAGTT	2,00			PTD013 protein
	GCAGGAGGTGA	2,00			chromosome 1 open reading frame 8
	GGAGCTGCTGC	2,00			Homo sapiens clone 25007 mRNA sequence
5051	GCCCGCCCTC	5,00	0,53	Hs.280666	Homo sapiens chromosome 19, cosmid R32184
5052	GAGGGAGTTTC	107,00	2,61	Hs.76064	ribosomal protein L27a
5053	TTCCTCCACGC	3,00			src homology 3 domain-containing protein HIP-55
5054	GCCTGGGCTGG	7,00	0.63	Hs.112184	DKFZP586J0619 protein
	GACAGTGTGGG	4,00			nuclear mitotic apparatus protein 1
	AATCCAGGAGG	5,00			dolichyl-diphosphooligosaccharide- protein glycosyltr
5057	CCCCTGGATC	31,00	1,30		S100 calcium-binding protein A6 (calcyclin)
5058	TACAGTATGTT	7,00	0,64		glutamate-ammonia ligase (glutamine synthase)
5059	CCCATCATCCC	8,00	0.68	Hs.109051	glycoprotein, synaptic 2
	GAGGAGGGTGA	4,00	0.48	Hs.75318	tubulin, alpha 1 (testis specific)
	GTCACAGTCCT	3,00			serum response factor (c-fos serum
					response element-
	GATGCCCTCCT	3,00			insulin receptor substrate 2
	GCCAGACCCCT	3,00			KIAA0515 protein
	TGCCTGTGGCC	2,00		Hs.7678	cellular retinoic acid-binding protein 1
5065	GGAAGAGCACT	2,00	0,30	Hs.75268	sialyltransferase 4C (beta-galactosidase alpha-2,3-s
F-0-0-0	CCCTCTGTGAT	2,00	0.30	Hs.74649	cytochrome c oxidase subunit VIc

<u> </u>	<u> </u>	0.00	0.00	11. 57704	l
5067	ATTGGCTGGGC	2,00	0,30	Hs.57764	protein phosphatase 1A (formerly 2C),
5000	COATTOACTOT	0.00	0.00	11- 404554	magnesium-depe
	GCATTGAGTGT	2,00			hypothetical protein FLJ20159
12028	TTTCCACTTAA	2,00	0,30	MS.101813	solute carrier family 9 (sodium/hydrogen
5070	00004047004	4.00	0.40	11- 00057	exchanger),
	CGGCACATCCA	1,00		Hs.92357	galactokinase 1
	ATACCAGATAC	1,00		Hs.9071	progesterone membrane binding protein
	AAGCACCTTGA	1,00		Hs.89862	TNFRSF1A-associated via death domain
	GGGGTCCCAAA	1,00		Hs.6815	ESTs
5074	TGTGAAAATAA	1,00	0,16	Hs.62349	ras association (RalGDS/AF-6) domain
					containing prot
	TCAAGAAATAA	1,00		Hs.61418	microfibrillar-associated protein 1
5076	CTGGTACCTGT	1,00	0,16	Hs.5813	ESTs, Moderately similar to 2109260A B cell growth f
5077	ACTCATCTGAC	1,00	0,16	Hs.5320	hypothetical protein
5078	AGGTATGGAGA	1,00	0,16	Hs.43654	hypothetical protein FLJ20561
	GGCACACCTTA	1,00	_	Hs.41127	Homo sapiens cDNA FLJ10745 fis, clone
					NT2RP3001676
5080	ATTGTAGACAA	1,00	0,16	Hs.39871	KIAA0727 protein
5081	GTCATATTTCC	1,00		Hs.32990	DKFZP566F084 protein
	TTGGAGGAGAG	1,00		Hs.32978	proprotein convertase subtilisin/kexin
		· 1	, i		type 7
5083	AGAGCAGGTAC	1,00	0,16		ESTs, Weakly similar to P137_HUMAN
			·		GPI-ANCHORED PROT
5084	TGGGTCTGGAT	1,00	0,16	Hs.25328	ESTs
5085	TGGAGAGAATA	1,00	0,16	Hs.249247	heterogeneous nuclear protein similar to rat helix d
5086	CCCAGGAGCAG	1,00	0.16	He 22051	Homo sapiens mRNA; cDNA
	COCACOACOAC	1,00	0,10	113.22001	DKFZp434O119 (from clone DKF
5087	ACACTAAGACG	1,00	0.16	Hs 170253	ESTs, Weakly similar to rabaptin-4
000,	710710171107100	1,00	0,10		[H.sapiens]
5088	GTCAGAACACC	1,00	0.16		zinc finger protein 42 (myeloid-specific
		1,00	0,10	1.0.100002	retinoic ac
5089	GTATAAACGAA	1,00	0.16	Hs.166982	phosphatidylinositol glycan, class F
	CCCCCTCCTTA	1,00			Homo sapiens mRNA; cDNA
		.,,	-,		DKFZp564D016 (from clone DKF
5091	GAAATGTATGC	1,00	0.16	Hs.142939	
	TCCTCTCACCA	1,00	0.16	Hs.12151	hypothetical protein FLJ11286
	GTGCTACTTCT	1,00			collagen, type IV, alpha 1
-	AATTCCAACTC	1,00			CGI-43 protein
	AGCTCCCAGAT	1,00			ferritin, light polypeptide
	TGTGAGGGCAT	1,00			hypothetical protein FLJ20602
	TTCCCTGTGTA	1,00			glucocorticoid receptor DNA binding
		',55	٥,.٠		factor 1
5098	TGCCTCTGCGG	18,00	1.01	Hs.75564	CD151 antigen
	GTAAAAAAAA	21,00			KIAA0242 protein
	GAACGCCTAAT	8,00			dihydropyrimidinase-like 2
	TCAGACGCAGC	40,00			prothymosin, alpha (gene sequence 28)
	AGCTGTCTCAA	3,00		Hs.99766	Homo sapiens mRNA; cDNA
					Transportation and America

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			<del></del> -	<del></del>	DVEZ-EGA 10202 /from clone DV
			0.40		DKFZp564J0323 (from clone DK
5103	ACGACAAAGCT	3,00	0,40	Hs.83920	peptidylglycine alpha-amidating
					monooxygenase
	CCCTTCTGCCA	3,00		Hs.6214	KIAA0731 protein
5105	ATAGCTGGGGC	3,00		Hs.3446	mitogen-activated protein kinase kinase 1
5106	GTTTCAGGAGT	3,00	0,40	Hs.156114	protein tyrosine phosphatase, non- receptor type subs
5107	CCTGTGTGCAT	3,00	0.40	Hs.11611	KIAA1424 protein
	CCCCAATGCT	3,00			splicing factor 3a, subunit 2, 66kD
	GGACCACTGAA	86,00			ribosomal protein L3
	TTTAAAAAAAA	2,00		Hs.74088	early growth response 3
	GGGCAGGGGAA	2,00			hypothetical protein FLJ20248
	TTGAGAGATGA	2,00		Hs.3758	COP9 complex subunit 7a
	AAGCAAAAGGT	2,00		Hs.28505	ubiquitin-conjugating enzyme E2H (homologous to yeas
5114	TGAAAAGCTTA	2,00	0,31	Hs.2384	tumor protein D52
5115	GCTCTGTTCAT	2,00	0,31	Hs.18192	Ser/Arg-related nuclear matrix protein (plenty of pr
5116	TTGAAGTGGTT	2,00	0,31	Hs.179972	hypothetical protein FLJ10154
5117	CGGAGCCGGCT	2,00	0,31	Hs.117582	CGI-43 protein
5118	GTCTGACCCCA	6,00	0,60	Hs.173902	protein phosphatase 2 (formerly 2A), regulatory subu
5119	CGACCCCACGC	30,00	1,35	Hs.169401	apolipoprotein E
	TACATAATTAC	7,00		Hs.120980	(Manual assignment) ORF-less transcript in MEN1 regi
5121	GCTGAAGATGA	3,00	0,41	Hs.78202	SWI/SNF related, matrix associated, actin dependent
5122	TGCTGTGCATA	8,00	0,70	Hs.147916	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 3
5123	CAGCCCAACCG	7,00	0,65	Hs.28081	eukaryotic translation initiation factor 3, subunit
5124	CCCTGGGTTCT	54,00	1,92	Hs.111334	ferritin, light polypeptide
5125	CTGTGGCCGGA	3,00	0,41	Hs.284296	Homo sapiens SURF-4 mRNA, complete cds
5126	CCTGTAAAGCC	2,00	0,31	Hs.9691	Homo sapiens cDNA FLJ11255 fis, clone PLACE1008902
5127	GCTTTTCCTGT	2,00	0,31	Hs.75277	Homo sapiens mRNA; cDNA DKFZp586M141 (from clone DKF
5128	ACCCTGGGCAC	2,00	0,31	Hs.220324	Human DNA sequence from clone 283E3 on chromosome 1p
5129	GCAGAGCCTTG	2,00	0,31	Hs.20768	HSPC189 protein
5130	AATATCTGACT	2,00	0,31	Hs.118021	active BCR-related gene
5131	GACATCCTGTC	1,00		Hs.87627	(Manual assignment) MEMOREC SAS10
5132	ATTTAGACCAG	1,00		Hs.84790	KIAA0225 protein
	TGCTGTGACCA	1,00		Hs.78465	v-jun avian sarcoma virus 17 oncogene homolog
5134	ATCGTGGAGGA	1,00	0,16	Hs.727	inhibin, beta A (activin A, activin AB alpha polypep

5135	TATTCAATTAC	1,00		Hs.7045	GL004 protein
5136	TCTTTGTCTAA	1,00	0,16	Hs.6838	ESTs
5137	GCCAAAGTGTT	1,00		Hs.5831	tissue inhibitor of metalloproteinase 1
			·		(erythroid p
5138	GATCCAAATGT	1,00	0,16	Hs.42650	ZW10 interactor
	GAGGCTTAATA	1,00			hypothetical protein
	AGGGGAGAGGA	1,00			trinucleotide repeat containing 15
	AGAGAAAAAA	1,00			Human XIST, coding sequence "a"
		.,	-,		mRNA (locus DXS399E)
5142	GATGTTGTCCA	1,00	0.16	Hs.279607	calpastatin
	CCCAATACTCT	1,00			hypothetical protein
	GAAATTGGTCT	1,00			heterogeneous nuclear protein similar to
		.,,,,	0,.0		rat helix d
5145	AACGAGTATTC	1,00	0.16	Hs.241578	U6 snRNA-associated Sm-like protein
		.,	٥,٠٠		LSm8
5146	TGCCAAAAAA	1,00	0.16		dystonia 1, torsion (autosomal dominant;
		.,,,,	٥,		torsin A)
5147	GGTGCACCCGG	1,00	0.16		Homo sapiens mRNA; cDNA
		.,,,,,	٠, . ٠		DKFZp434N1221 (from clone DK
5148	TCTAGCATTTC	1,00	0.16	Hs 179260	ESTs, Moderately similar to T12543
		.,00	0, 10		hypothetical prot
5149	AGCTTTGTAGA	1,00	0.16		hypothetical protein FLJ10955
	CCCAAGGTCTT	1,00		Hs.153818	
-	AACCAATACAG	1,00	0,10	He 146161	ESTs, Weakly similar to KIAA0859
0.0.		1,00	0,10		protein [H.sapiens]
5152	CTTTTTAAATC	1,00	0.16		suppression of tumorigenicity 13 (colon
0.02		1,00	0, 10	113.110222	carcinoma) (
5153	GCTCCTGAGCC	1,00	0.16	He 111088	PR/SET domain containing protein 07
	GAGAAACATTT	1,00		Hs.101619	
	CAAAATGCAAA	1,00			cathepsin C
	TGGAATGCTGG	12,00			NADH dehydrogenase (ubiquinone)
3 130	IGOANIGCIGG	12,00	0,07	П5.//44	flavoprotein 1 (51kD
5157	ACCTTGTGCCC	5,00	0.55		sorbitol dehydrogenase
	TGGAGTGGAGG	17,00			guanylate kinase 1
	CTCCACAAATT	4,00			PDGF associated protein
	AGAAATCACTG				
5100	AGAMATCACTO	2,00	0,31		L-3-hydroxyacyl-Coenzyme A
5161	TGCAGGTGTGT	2 00	0.24		dehydrogenase, short chai
		2,00			high-glucose-regulated protein 8
3 102	CCTGCAGTCCC	2,00	0,31		ESTs, Weakly similar to ALU8_HUMAN
E160	CCCAAACTTTC	2.00	0.34		ALU SUBFAMILY SX
اكفا دا	CCCAAACTTTG	2,00	0,31	ms.109438	Homo sapiens clone 24775 mRNA
5164	CCCTCCAAAAA	2 00	0.40	H- 040405	sequence
2 104	GGGTGCAAAAA	3,00	0,42	⊓5.∠49495	heterogeneous nuclear ribonucleoprotein
E405	CTATCACTUTE		0.40	11- 40 4005	A1
2100	CTATCAGTTTT	3,00	0,42	ms.194625	dynein, cytoplasmic, light intermediate
E400	ATTOTOGAGEA	70.00	0.46	11- 004540	polypeptide
	ATTCTCCAGTA	73,00			ribosomal protein L23
	GCAAAGAAAAA	4,00			LIM domain only 4
5168	ATGAGCTGACC	7,00	0,68	Hs.695	cystatin B (stefin B)

5169	TAATTTGCATT	3,00			epithelial membrane protein 1
5170	AAACACTCTTG	3,00	0,42		oxidase (cytochrome c) assembly 1-like
5171	CCAACAAGAAT	4,00	0,50	Hs.82749	transmembrane 4 superfamily member 2
5172	TTAAAGGCCGG	4,00	0,50	Hs.79086	ribosomal protein, mitochondrial, L3
5173	GCCGAGACCAA	2,00	0,32	Hs.61258	argininosuccinate lyase
5174	GTTTAAGTTAA	2,00	0,32	Hs.30029	ESTs
	CCTTGCTTTTA	2,00	0,32	Hs.278736	cell division cycle 42 (GTP-binding
					protein, 25kD)
5176	TGAGGACACAG	2,00	0,32	Hs.14541	cullin 1
5177	TTGGGAATCCC	1,00	0,16	Hs.9547	hypothetical protein FLJ10916
5178	CATTTTTCCCC	1,00	0,16	Hs.90336	ATPase, H+ transporting, lysosomal
				•	(vacuolar proton
5179	ACTCAAATCTT	1,00	0,16	Hs.7953	HSPC041 protein
	TAATTTTAACT	1,00		Hs.78867	protein tyrosine phosphatase, receptor-
					type, Z polyp
5181	CAGGTGTCTTT	1,00	0,16	Hs.77100	general transcription factor IIE,
					polypeptide 2 (bet
5182	ACTGGTGAGAG	1,00	0,16	Hs.76728	ESTs
	AAAATAAAGCT	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	ACTCACTGCAG	1,00	0,16	Hs.60377	KIAA1298 protein
	TTAAAATTGCC	1,00	0.16	Hs.444	serine/threonine kinase 19
	AACCAGGTGGA	1,00		Hs.44095	hypothetical protein FLJ20018
	TAGGTCAGGAC	1,00		Hs.43666	protein tyrosine phosphatase type IVA,
		,,,,,	.,		member 3
5188	GTGGCGGCACC	1,00	0,16	Hs.43112	Homo sapiens mRNA; cDNA
					DKFZp434B1620 (from clone DK
5189	CCCACAACCCG	1,00	0,16	Hs.38205	from HeLa cyclin-dependent kinase 2
					interacting prot
5190	GAATTGAGCTT	1,00	0,16	Hs.36787	chromodomain helicase DNA binding
					protein 2
	TGTTCTTTGCA	1,00		Hs.3376	hypothetical protein FLJ10743
5192	GCTTCCTCTGC	1,00	0,16	Hs.27262	Homo sapiens clone 25110 mRNA
					sequence
5193	GGATACAACAC	1,00	0,16	Hs.272458	protein phosphatase 3 (formerly 2B),
					catalytic subun
5194	GTAGTCACCGC	1,00	0,16	Hs.26630	ATP-binding cassette, sub-family A
					(ABC1), member 3
	GCACAAGAGTG	1,00		Hs.25892	ESTs
	GCCTCCAGCCT	1,00		Hs.23100	ESTs
	TAAACATTGTC	1,00		Hs.23060	DKFZP564F0522 protein
5198	CAAATTACAAT	1,00		Hs.22393	density-regulated protein
5199	CTTACAGCCAC	1,00		Hs.195969	
5200	ACTGCACCACT	1,00	0,16	Hs.185910	ESTs, Weakly similar to ALU1_HUMAN
L					ALU SUBFAMILY J S
5201	TTACAATGCTG	1,00	0,16	Hs.184013	ESTs, Highly similar to unnamed protein
L					product [H.s
5202	TCCTACGGAAA	1,00	0,16	Hs.108708	calcium/calmodulin-dependent protein
					kinase kinase 2
5203	GCTGTTCATTG	3,00	0,42	Hs.77306	survival of motor neuron 1, telomeric

5204	ATATAATCTGA	3,00	0,42	Hs.621	lectin, galactoside-binding, soluble, 3
					(galectin 3)
$\overline{}$	CTAATAAACTT	3,00			CGI-81 protein
5206	CAGGATCCAGA	6,00	0,63	Hs.119222	suppression of tumorigenicity 13 (colon carcinoma) (
5207	CGGATAACCAG	7,00	0,69	Hs.5181	proliferation-associated 2G4, 38kD
5208	GTGTATCTTTT	5,00	0,57	Hs.73965	splicing factor, arginine/serine-rich 2
5209	CTCCACCTGGT	2,00	0,32	Hs.75835	phosphomannomutase 1
5210	TCAACTGGTTC	2,00	0,32	Hs.75812	phosphoenolpyruvate carboxykinase 2 (mitochondrial)
5211	AAATTTTAAAA	2,00	0,32	Hs.74649	cytochrome c oxidase subunit VIc
5212	ACCCACAGTGC	2,00	0,32	Hs.67896	7-60 protein
5213	AGGCCTGGGCC	2,00	0,32	Hs.6163	ESTs, Weakly similar to myotonic dystrophy kinase [H
5214	TCAGTGCGCAG	2,00	0,32	Hs.5534	ESTs
5215	CTGAGGTGATG	2,00	0,32	Hs.26367	Homo sapiens mRNA; cDNA
					DKFZp564M1178 (from clone DK
	CCACTCCACTC	2,00	0,32	Hs.190452	KIAA0365 gene product
5217	GCCTGCTCCCT	2,00	0,32	Hs.16725	hypothetical protein FLJ10111
5218	GACTGTGCCAC	11,00	0,88	Hs.5120	dynein, cytoplasmic, light polypeptide
5219	TGATCTCCAAA	8,00	0,74	Hs.83190	fatty acid synthase
5220	AAGAAAGTTCT	5,00	0,58	Hs.10729	hypothetical protein
5221	TAACAGAAAGG	3,00	0,43	Hs.97496	YY1 transcription factor
5222	ATGGCTAAGCT	3,00	0,43	Hs.82280	regulator of G-protein signalling 10
5223	GATGGGGACAA	3,00	0,43	Hs.118724	DR1-associated protein 1 (negative cofactor 2 alpha)
5224	AATTCAATTAA	4,00	0,51	Hs.211568	eukaryotic translation initiation factor 4 gamma, 1
5225	AGCAAACTGAA	4,00	0,51	Hs.182579	leucine aminopeptidase
5226	CCTCGGAAAAT	30,00			ribosomal protein L38
	ATGGCCCATAC	2,00			carboxylesterase 2 (intestine, liver)
5228	CACCACGGGCC	2,00			breast cancer anti-estrogen resistance 1
5229	CGGATAAGGCC	2,00			nuclear prelamin A recognition factor
5230	CCTTGGGCCTA	2,00			chromatin-specific transcription elongation factor,
5231	TATAACTTGTA	2,00	0,32	Hs.132955	BCL2/adenovirus E1B 19kD-interacting protein 3-like
5232	GCACCTTATTG	2,00	0,32	Hs.125078	ornithine decarboxylase antizyme 1
	CACATCTCTGA	2,00			Homo sapiens cDNA FLJ20167 fis, clone COL09512
5234	CCTTGTCCAGC	2,00	0,32	Hs.101067	GCN5 (general control of amino-acid synthesis, yeast
5235	AGCTGGTTTCC	6,00	0,65	Hs.286027	etoposide-induced mRNA
5236	TTCCCTCGTGA	3,00	0,43	Hs.80758	aspartyl-tRNA synthetase
5237	CGCACCATTGC	5,00	0,58	Hs.94672	GCN5 (general control of amino-acid
E000	ATATTOTOGOT		0.4=	11-0000	synthesis, yeast
	ATATTCTGCCT	1,00			hypothetical protein
	CTCCCAAGCTC	1,00		Hs.9452	KIAA0770 protein
5240	CAAGCTTGGTC	1,00	υ,17	Hs.86858	ribosomal protein S6 kinase, 70kD,

	,	,			
					polypeptide 1
	GTCATCACTGG	1,00		Hs.8535	hypothetical protein bA395L14.2
5242	CCCTGTTCAGC	1,00	0,17	Hs.78824	tyrosine kinase with immunoglobulin and
<u> </u>					epidermal gr
5243	GACTGGAAAAA	1,00	0,17	Hs.743	Fc fragment of IgE, high affinity I,
					receptor for; g
5244	GACCCTTCTCC	1,00	0,17	Hs.6580	Homo sapiens clone 23718 mRNA
					sequence
5245	GGCAGGAGTAG	1,00	0,17	Hs.62661	guanylate binding protein 1, interferon-
					inducible, 6
5246	TGCTAGGAAGG	1,00	0,17	Hs.55235	sphingomyelin phosphodiesterase 2,
				Υ.	neutral membrane
5247	TGTTGTATTTG	1,00	0,17		ESTs
5248	GTGGCACCCGC	1,00	0,17	Hs.47305	ESTs, Weakly similar to LONN_HUMAN
		, l	·		MITOCHONDRIAL LON
5249	AAACTCACGCC	1,00	0,17	Hs.3337	transmembrane 4 superfamily member 1
	TGTATGGCTGG	1,00			Homo sapiens calpain 3 (CAPN3)
			•		mRNA, complete cds, a
5251	GCTTAAATTAA	1,00	0.17	Hs.282283	baculoviral IAP repeat-containing 2
	AGAAGATTTAT	1,00		Hs.250746	
	ACCTGGCCTGA	1,00			tuftelin-interacting protein
	CTGTACATACT	1,00			NS1-binding protein
	ATGCTGGGGAG	1,00			Homo sapiens mRNA for P53TG1-D,
00		',55	٥,		complete cds
5256	GCAATAATGGT	1,00	0.17	Hs.19614	
	GCTGGCCGGAA	1,00			GTPase activating protein-like
	CGCTGGTTCCC	1,00			ribosomal protein L11
	CAGGGGCTGGG	1,00			KIAA0677 gene product
	CGTTTAATCAT	1,00			E1B-55kDa-associated protein 5
	CTCCCAGGTCA	1,00			M-phase phosphoprotein 6
	AGCCCTCAACA	1,00			DKFZP564F1422 protein
_	GACTGAATGTA	1,00			ESTs, Weakly similar to alternatively
5203	GACIGAAIGIA	1,00	0, 17		spliced produc
5264	GTAATGCATAT	1,00	0.17	Un 4004E0	SH3-domain binding protein 5 (BTK-
5204	JANIGCAIAI	1,00	0,17	ms. 109 150	associated)
E265	CTCACCCCACA	1.00	0.17	Un 107292	KIAA1517 protein
	GTGAGGGCACA	1,00			
<b>5200</b>	CCCTGGCAATG	5,00	0,59	MS.273369	uncharacterized hematopoietic
5067	ACCCACCACA	52.00	0.00	11- 400500	stem/progenitor cells
	ACGCAGGAGA	53,00			heat shock 90kD protein 1, alpha
	CCCGGGAGCGA	4,00			carboxy terminal LIM domain protein 1
0269	GAGGCCATCCC	4,00	0,52	Hs.70830	U6 snRNA-associated Sm-like protein
	0007007070			11. 00405	LSm7
52/0	GGCTGGTCTCC	6,00	0,65	Hs.86185	Alu-binding protein with zinc finger
	0400000				domain
	GAGGGCCTTGT	2,00			tuberous sclerosis 2
	GTGACAGAATT	2,00			UDP-glucose pyrophosphorylase 2
	CTTTCACTTC	2,00		Hs.279919	
5274	TGTGTTGTGTC	2,00	0,33	Hs.279806	Homo sapiens mRNA; cDNA
L	<u> </u>	łl			DKFZp434E109 (from clone DKF

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52/5	TGGGGTGGAGT	2,00	0,33	Hs.26403	glutathione transferase zeta 1
5070	0470440044	0.00	0.00	11 400040	(maleylacetoacetate i
52/6	GATTCAACCAA	2,00	0,33	Hs.168213	ESTs, Weakly similar to ALU1_HUMAN
					ALU SUBFAMILY J S
	CCGTCATCCTG	2,00			Not56 (D. melanogaster)-like protein
-	AGCTGAGCTAA	2,00			deoxyribonuclease II, lysosomal
	CAATAAATGTT	48,00			ribosomal protein L37
5280	GCAGGGCCTCA	13,00	1,01	Hs.92323	FXYD domain-containing ion transport
					regulator 3
5281	CAGTTACTTAG	9,00	0,83	Hs.279920	tyrosine 3-monooxygenase/tryptophan 5-
					monooxygenase
5282	TTCCTGGTAGT	6,00			KIAA0788 protein
5283	GTTTTCATTCA	5,00	0,60	Hs.173736	ancient ubiquitous protein 1
5284	CCCTCCTCTCC	2,00	0,33	Hs.83173	cyclin D3
5285	TCCCTGGCAGA	2,00	0,33	Hs.70327	cysteine-rich protein 2
5286	GACACGAACAA	2,00	0,33		ras-related protein
5287	CTGGGATGTCG	2,00	0.33		CGI-92 protein
	GTGTTCTGACT	2,00	0.33		DKFZP586P2220 protein
	GCTATGCTCCC	2,00			hypothetical protein PRO2207
	GCCCTCAGCA	2,00			hypothetical protein FLJ20419
	CCCCCTGCCCT	7,00			hypothetical protein FLJ10350
	AGAAAGATGTC	11,00			annexin A1
	TTTACAGCTGG	3,00		Hs.89981	diacylglycerol kinase, zeta (104kD)
	GTTGTAAAATA	3,00		Hs.7869	lysophosphatidic acid acyltransferase-
3294	GIIGIAWAIA	3,00			delta
5295	CTGCCTCCTTA	4,00	0,53	Hs.7918	uncharacterized hypothalamus protein HSMNP1
	TCACCTTCAAG	1,00	0,17	Hs.74002	nuclear receptor coactivator 1
5297	TTTTATAAGGA	1,00	0,17	Hs.73986	CDC-like kinase 2
5298	CCTTGTCCTCT	1,00	0,17	Hs.69743	GM2 ganglioside activator protein
5299	AGTGTTTGTAG	1,00	0,17	Hs.5420	hypothetical protein FLJ20695
5300	AGAACAAAGGC	1,00	0,17	Hs.50785	SEC22, vesicle trafficking protein (S. cerevisiae)-l
5301	GGCAAAAAAA	1,00	0,17	Hs.2953	ribosomal protein S15a
	GCCTCCACAGC	1,00		Hs.285813	
=	GCACAAGTTCT	1,00			hypothetical protein FLJ20216
	GTTGGGTAGAA	1,00			Human DNA sequence from clone RP1-
			·		28H20 on chromosom
5305	AGGTCAGGAAA	1,00	0.17	Hs.249429	
	GTCATTTTCTA	1,00			transcription factor 8 (represses
		.,	-,		interleukin 2 expr
5307	TCTTCCCTCAG	1,00	0.17		ESTs, Moderately similar to
		.,	-,		ALU7_HUMAN ALU SUBFAMILY
5308	GGTTGATCACC	1,00	0.17	Hs.19699	Conserved gene telomeric to alpha
		.,	-,		globin cluster
5309	TTTCAAGTGGT	1,00	0 17	Hs.17820	Rho-associated, coiled-coil containing
		.,55	٥,		protein kinas
5310	CCGGGTTATTT	1,00	0 17	Hs 160138	RPA-binding trans-activator
	CCGCCCCCAGC	1,00			islet cell autoantigen 1 (69kD)
0011	COGCOCCAGO	ןטטן, ו	0,17	113.10/82/	polet cell autoantigen 1 (03KD)

S312   GTGTTGACTGC	5313 TGAAACGGAAG         1,00         0,17 Hs.12940         zinc-fingers and homeobted standard of the control of the con	oin
S314   GTGCTCAGCCT	S314   GTGCTCAGCCT	
S315 ACTGGTGTCT	Display	
S315   ACCTGGTGTCT   1,00   0,17   Hs. 111988   PR/SET domain containing protein 07   5316   GCAGAGAAAAA   1,00   0,17   Hs. 109606   coronin, actin-binding protein, 1A   5317   TAAGTGACTGT   1,00   0,17   Hs. 103606   coronin, actin-binding protein, 1A   5318   CACAGAGTCCT   4,00   0,53   Hs. 75140   low density lipoprotein-related protein-associated p   low density lipoprotein   low density	5315 ACCTGGTGTCT         1,00         0,17 Hs.111988 PR/SET domain containing           5316 GCAGAGAAAAA         1,00         0,17 Hs.109606 coronin, actin-binding pro           5317 TAAGTGACTGT         1,00         0,17 Hs.103755 receptor-interacting serine kinase 2           5318 CACAGAGTCCT         4,00         0,53 Hs.75140 low density lipoprotein-rel associated p           5319 TAAACTGTTTC         9,00         0,85 Hs.3491 ribosomal protein S14           5320 TTCACAGATTT         3,00         0,44 Hs.8107 Homo sapiens mRNA; cD DKFZp58680918 (from closs)           5321 AGCTGATCAGC         3,00         0,44 Hs.78223 N-acylaminoacyl-peptide           5322 TTCATTGTAGA         4,00         0,53 Hs.23960 cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.284271 cytochrome b5 reductase           5325 CACTGTGACCT         2,00         0,33 Hs.15829 KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10882 HMG-box containing protein           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550 myosin, heavy polypeptide muscle           5331 GCCAGACACCC         3,00         0,45 Hs.89781 upstream binding transcriperate protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730 lheterogeneous nuclear rise protein           5333 TAGCCAGACACCC         3,00         0,45 Hs.89801 livB (bacterial acetolactate protein protein	ınnamed
5316         GCAGAGAAAAA         1,00         0,17         Hs.109765         coronin, actin-binding protein, 1A           5317         TAAGTGACTGT         1,00         0,17         Hs.103755         receptor-Interacting serine-threonine kinase 2           5318         CACAGAGTCCT         4,00         0,53         Hs.75140         low density lipoprotein-related protein-associated p           5319         TTAAACTGTTTC         9,00         0,85         Hs.3491         ribosomal protein S14           5320         TTCACAGATTT         3,00         0,44         Hs.8107         Homo sapiens mRNA; cDNA DKFZp58680918 (from clone DK           5321         AGCTGATCAGC         3,00         0,44         Hs.78223         N-acylaminoacyl-peptide hydrolase           5322         TTCATTGTAGA         4,00         0,53         Hs.6527         G protein-coupled receptor 56           5323         TGCATCTGTA         4,00         0,53         Hs.23960         cyclin B1           5326         CACTGTGACCT         2,00         0,33         Hs.23960         cyclin B1           5327         CACTGTGACCT         2,00         0,33         Hs.15829         KlAA0676 protein           5326         CACTGTGACCT         2,00         0,33         Hs.10443         Homo sapiens mRNA; cDN	5316         GCAGAGAAAAA         1,00         0,17         Hs. 109606         coronin, actin-binding prosental forms of the protein o	
5317 TAAGTGACTGT         1,00         0,17 Hs.103755 receptor-interacting serine-threonine kinase 2           5318 CACAGAGTCCT         4,00         0,53 Hs.75140 low density lipoprotein-related protein-associated p           5319 TAAACTGTTTC         9,00         0,85 Hs.3491 ribosomal protein S14           5320 TTCACAGATTT         3,00         0,44 Hs.8107 low osapiens mRNA; cDNA DKFZp58680918 (from clone DK           5321 AGCTGATCAGC         3,00         0,44 Hs.78223 low osapiens mRNA; cDNA DKFZp58680918 (from clone DK           5322 TTCATTGTAGA         4,00         0,53 Hs.5527 loy osapiens mRNA; cDNA DKFZp58680918 (from clone DK           5322 GCATCTGTA         4,00         0,53 Hs.29860 loy osapiens mRNA; cDNA DKFZp58680918 (from clone DKF           5325 GACTGTGACCT         2,00         0,33 Hs.7765 low chrome b5 reductase 1 (B5R.1)           5326 GGCTGCTCTT         2,00         0,33 Hs.155829 kIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10882 lhMG-box containing protein 1           5328 ATGTTGCCCT         2,00         0,33 Hs.10882 lhMG-box containing protein 1           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550 myosin, heavy polypeptide 9, non-muscle           5331 GCCAGACACCC         3,00         0,45 Hs.89781 low polymeras           5332 TACCTTTGGCA         14,00         1,11 Hs.2730 low polymeras           5333 TAGGCCCAAGT	5317 TAAGTGACTGT         1,00         0,17 Hs.103755         receptor-interacting serind kinase 2           5318 CACAGAGTCCT         4,00         0,53 Hs.75140         low density lipoprotein-rel associated p           5319 TAAACTGTTTC         9,00         0,85 Hs.3491         ribosomal protein S14           5320 TTCACAGATTT         3,00         0,44 Hs.8107         Homo sapiens mRNA; cD DKFZp586B0918 (from cl DKFZp58B0 (from cl DKFZp58B0) (from cl DKFZp58B0 (from cl DKFZp58B0) (from cl DKFZp58B0 (from cl DKFZp58B0) (from cl DKFZp58B0) (from cl DKFZp58B0 (from cl DKFZp58B0) (from cl DKFZp58B0) (from cl DKFZp58B0 (from cl DKFZp58B0)	
Single Cacagactic	Sala   CACAGAGTCCT   4,00   0,53   Hs.75140   low density lipoprotein-relassociated p   associated p   associated p   associated p   associated p   associated p   Tibosomal protein S14   Sala   Transcription   Transcript	
S318   CACAGAGTCCT	5318         CACAGAGTCCT         4,00         0,53         Hs.75140         low density lipoprotein-rel associated p           5319         TAAACTGTTTC         9,00         0,85         Hs.3491         ribosomal protein S14           5320         TTCACAGATTT         3,00         0,44         Hs.8107         Homo sapiens mRNA; cD DKFZp586B0918 (from clooper per per per per per per per per per	e-threonine
S319   TAAACTGTTTC	Sassociated p   Sassociated	
S319   TAAACTGTTTC	5319 TAAACTGTTTC         9,00         0,85 Hs.3491         ribosomal protein S14           5320 TTCACAGATTT         3,00         0,44 Hs.8107         Homo sapiens mRNA; cD DKFZp586B0918 (from close)           5321 AGCTGATCAGC         3,00         0,44 Hs.78223         N-acylaminoacyl-peptide           5322 TTCATTGTAGA         4,00         0,53 Hs.6527         G protein-coupled receptor           5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.7765         chromosome 16 open real chromosome 15 reductase           5326 GGGCTGCTCTT         2,00         0,33 Hs.155829 KlAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10882         HMG-box containing protein chromosome lange protein chromosome lange protein chromosome lange protein chromosome lange protein lan	lated protein-
S320   TTCACAGATTT   3,00	5320         TTCACAGATTT         3,00         0,44         Hs.8107         Homo sapiens mRNA; cD DKFZp586B0918 (from cl DKFZp586B0918)           5321         AGCTGATCAGC         3,00         0,44         Hs.78223         N-acylaminoacyl-peptide           5322         TTCATTGTAGA         4,00         0,53         Hs.6527         G protein-coupled receptor           5323         TGCCATCTGTA         4,00         0,53         Hs.23960         cyclin B1           5324         GCCTCCTGTCA         2,00         0,33         Hs.7765         chromosome 16 open real chro	
DKFZpS86B0918 (from clone DK	DKFZp586B0918 (from cl   5321 AGCTGATCAGC   3,00   0,44 Hs.78223   N-acylaminoacyl-peptide   5322 TTCATTGTAGA   4,00   0,53 Hs.6527   G protein-coupled receptor   5323 TGCCATCTGTA   4,00   0,53 Hs.23960   cyclin B1   5324 GCCTCCTGTCA   2,00   0,33 Hs.7765   chromosome 16 open rea   5325 CACTGTGACCT   2,00   0,33 Hs.155829 KIAA0676 protein   5327 CCCAGGACACC   2,00   0,33 Hs.155829 KIAA0676 protein   5328 ATGTTGCCCT   2,00   0,33 Hs.10882   HMG-box containing protein   5329 TGCTAAAAAAA   6,00   0,68 Hs.146550   myosin, heavy polypeptide   muscle   1,00   1,11 Hs.2730   heterogeneous nuclear rik   L   1,00   1,11 Hs.2730   heterogeneous nuclear rik   L   1,00   1,11 Hs.2730   heterogeneous nuclear rik   1,333 TAGGCCCAAGT   4,00   0,54 Hs.78880   ilvB (bacterial acetolactate   5334 CCAACCGTGCT   8,00   0,81 Hs.75207   glyoxalase   1,00   1,54 Hs.83623   nuclear receptor subfamily member 3   1,00   1	
5321 AGCTGATCAGC         3,00         0,44 Hs.78223         N-acylaminoacyl-peptide hydrolase           5322 TTCATTGTAGA         4,00         0,53 Hs.6527         G protein-coupled receptor 56           5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.7765         chromosome 16 open reading frame 5           5325 CACTGTGACCT         2,00         0,33 Hs.284271         cytochrome b5 reductase 1 (B5R.1)           5326 GGGCTGCTCTT         2,00         0,33 Hs.155829         KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10842         HMG-box containing protein 1           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing protein 1           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptide 9, non-muscle           5331 GCCAGACACCC         3,00         0,45 Hs.89781         upstream binding transcription factor, RNA polymeras           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear ribonucleoprotein l.           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880         liVB (bacterial acetolactate synthase)-like           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACC	5321 AGCTGATCAGC         3,00         0,44 Hs.78223         N-acylaminoacyl-peptide           5322 TTCATTGTAGA         4,00         0,53 Hs.6527         G protein-coupled recepto           5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.7765         chromosome 16 open rea           5325 CACTGTGACCT         2,00         0,33 Hs.15829         KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10882         HMG-box containing protein           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing protein           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptid muscle           5330 GGGGAGGGGGC         3,00         0,45 Hs.89781         upstream binding transcrig RNA polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear ris           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfa	
5322 TTCATTGTAGA         4,00         0,53 Hs.6527         G protein-coupled receptor 56           5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.27765         chromosome 16 open reading frame 5           5325 CACTGTGACCT         2,00         0,33 Hs.284271         cytochrome b5 reductase 1 (B5R.1)           5326 GGGCTGCTCT         2,00         0,33 Hs.155829 KlAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10882 HMG-box containing protein 1           5328 ATGTTGCCCT         2,00         0,33 Hs.146550           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550           5330 GGGGAGGGGC         3,00         0,45 Hs.89781         upstream binding transcription factor, RNA polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333 TAGGCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate synthase)-like           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5336 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily 1, group I, member 3	5322         TTCATTGTAGA         4,00         0,53         Hs.6527         G protein-coupled receptor           5323         TGCCATCTGTA         4,00         0,53         Hs.23960         cyclin B1           5324         GCCTCCTGTCA         2,00         0,33         Hs.7765         chromosome 16 open rea           5325         CACTGTGACCT         2,00         0,33         Hs.155829         KIAA0676 protein           5327         CCCAGGACACC         2,00         0,33         Hs.110443         Homo sapiens mRNA; cD           5328         ATGTTGCCCCT         2,00         0,33         Hs.110443         Homo sapiens mRNA; cD           5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptid muscle           5330         GGGGAGGGGGC         3,00         0,45         Hs.89781         upstream binding transcrig RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear rit           L         L         S333         TAGGCCCAAGT         4,00         0,54         Hs.83623         invelage receptor subfamily member 3	
5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.23960         cyclin B1           5325 CACTGTGACCT         2,00         0,33 Hs.284271         cytochrome b5 reductase 1 (B5R.1)           5326 GGGCTGCTCTT         2,00         0,33 Hs.155829         KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10443         Homo sapiens mRNA; cDNA DKFZp7610051 (from clone DKF           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing protein 1           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptide 9, non-muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333 TAGGCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate synthase)-like           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5336 GCATTTGACAG         2,00         0,34 Hs.88862         nuclear receptor subfamily 1, group I, member 3           5337 GGTGCGGCTGG         2,00 <td>5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.7765         chromosome 16 open rea           5325 CACTGTGACCT         2,00         0,33 Hs.284271         cytochrome b5 reductase           5326 GGGCTGCTCTT         2,00         0,33 Hs.155829         KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10443         Homo sapiens mRNA; cD DKFZp7610051 (from closmostal containing protein)           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing protein           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptid muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781         upstream binding transcrigen was closmostal contained by polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear rist           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily member 3           5336 GCATTTGACAG</td> <td></td>	5323 TGCCATCTGTA         4,00         0,53 Hs.23960         cyclin B1           5324 GCCTCCTGTCA         2,00         0,33 Hs.7765         chromosome 16 open rea           5325 CACTGTGACCT         2,00         0,33 Hs.284271         cytochrome b5 reductase           5326 GGGCTGCTCTT         2,00         0,33 Hs.155829         KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.10443         Homo sapiens mRNA; cD DKFZp7610051 (from closmostal containing protein)           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing protein           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptid muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781         upstream binding transcrigen was closmostal contained by polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear rist           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily member 3           5336 GCATTTGACAG	
5324         GCCTCCTGTCA         2,00         0,33         Hs.7765         chromosome 16 open reading frame 5           5325         CACTGTGACCT         2,00         0,33         Hs.284271         cytochrome b5 reductase 1 (B5R.1)           5326         GGGCTGCTCTT         2,00         0,33         Hs.155829         KIAA0676 protein           5327         CCCAGGACACC         2,00         0,33         Hs.110443         Homo sapiens mRNA; cDNA DKFZP7610051 (from clone DKF           5328         ATGTTGCCCCT         2,00         0,33         Hs.10882         HMG-box containing protein 1           5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptide 9, nonmuscle           5331         GGGGAGGGGC         3,00         0,45         Hs.89781         upstream binding transcription factor, RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.83804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333         TAGGCCAAGT         4,00         0,54         Hs.78880         iivB (bacterial acetolactate synthase)-like           5334         CCAACCGTGCT         8,00         0,81	5324         GCCTCCTGTCA         2,00         0,33         Hs.7765         chromosome 16 open rea           5325         CACTGTGACCT         2,00         0,33         Hs.284271         cytochrome b5 reductase           5326         GGGCTGCTCTT         2,00         0,33         Hs.155829         KIAA0676 protein           5327         CCCAGGACACC         2,00         0,33         Hs.10443         Homo sapiens mRNA; cD DKFZp7610051 (from closure)           5328         ATGTTGCCCCT         2,00         0,33         Hs.10882         HMG-box containing protein           5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptid muscle           5330         GGGGAGGGGC         3,00         0,45         Hs.89781         upstream binding transcrigen with the polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear rit           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         ilvB (bacterial acetolactate           5334         CCAACCGTGCT         8,00         0,81         Hs.75207         glyoxalase I <t< td=""><td>or 56</td></t<>	or 56
5325         CACTGTGACCT         2,00         0,33         Hs.284271         cytochrome b5 reductase 1 (B5R.1)           5326         GGGCTGCTCTT         2,00         0,33         Hs.155829         KIAA0676 protein           5327         CCCAGGACACC         2,00         0,33         Hs.110443         Homo sapiens mRNA; cDNA           DKFZP7610051         (from clone DKF           5328         ATGTTGCCCCT         2,00         0,33         Hs.10882         HMG-box containing protein 1           5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptide 9, non-muscle           5330         GGGGAGGGGC         3,00         0,45         Hs.89781         upstream binding transcription factor, RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         iiVB (bacterial acetolactate synthase)-like           5334         CCAACCGTGCT         8,00         0,81         Hs.75207         glyoxalase I           5336         GCATTTGACAG	5325 CACTGTGACCT         2,00         0,33 Hs.284271 cytochrome b5 reductase           5326 GGGCTGCTCTT         2,00         0,33 Hs.155829 KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.110443 Homo sapiens mRNA; cD DKFZp7610051 (from closure) DKFZp7610051 (from clos	
5326 GGGCTGCTCTT         2,00         0,33 Hs.155829 KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.110443 Homo sapiens mRNA; cDNA DKFZp7610051 (from clone DKF           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882 HMG-box containing protein 1           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550 myosin, heavy polypeptide 9, non-muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781 upstream binding transcription factor, RNA polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804 DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730 heterogeneous nuclear ribonucleoprotein Lus           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880 livB (bacterial acetolactate synthase)-like           5334 CCAACCGTGCT         8,00         0,81 Hs.75207 glyoxalase I nuclear receptor subfamily 1, group I, member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649 cytochrome c oxidase subunit VIc           5337 GGTGCGGCTGG         2,00         0,34 Hs.188882 Homo sapiens clone 23872 mRNA sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.170980 ESTs, Weakly similar to CBF1 interacting corepressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318 HS1 binding protein           5341 GTGCCTAGGA         3,00         0,45 Hs.12854 ATRAP protein           5342 AGTATCTGG	5326 GGGCTGCTCTT         2,00         0,33 Hs.155829 KIAA0676 protein           5327 CCCAGGACACC         2,00         0,33 Hs.110443 Homo sapiens mRNA; cD DKFZp7610051 (from closure protein)           5328 ATGTTGCCCCT         2,00         0,33 Hs.10882 HMG-box containing protein           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550 myosin, heavy polypeptide muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781 upstream binding transcription muscle           5331 GCCAGACACCC         3,00         0,45 Hs.3804 DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730 heterogeneous nuclear rist           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880 ilvB (bacterial acetolactate protein)           5335 GGTGACCACCA         4,00         0,54 Hs.83623 nuclear receptor subfamily member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649 cytochrome c oxidase subsequence           5338 TTGTGATTAAT         2,00         0,34 Hs.18882 Homo sapiens clone 2387 sequence           5339 GGAGAGACAGG         2,00         0,34 Hs.18442 E-1 enzyme           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318 HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854 ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.12854 ATRAP protein	
5327         CCCAGGACACC         2,00         0,33         Hs.110443         Homo sapiens mRNA; cDNA DKFZp7610051 (from clone DKF           5328         ATGTTGCCCCT         2,00         0,33         Hs.10882         HMG-box containing protein 1           5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptide 9, non-muscle           5330         GGGGAGGGGC         3,00         0,45         Hs.89781         mystream binding transcription factor, RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         ilvB (bacterial acetolactate synthase)-like           5334         CCAACCGTGCT         8,00         0,81         Hs.75207         glyoxalase I           5335         GGTGACCACCA         4,00         0,54         Hs.83623         nuclear receptor subfamily 1, group I, member 3           5336         GCATTTGACAG         2,00         0,34         Hs.188882         Homo sapiens clone 23872 mRNA sequence           5338         TTGGATTAAT         2,00         <	5327         CCCAGGACACC         2,00         0,33         Hs.110443         Homo sapiens mRNA; cD DKFZp7610051 (from closed possible)           5328         ATGTTGCCCCT         2,00         0,33         Hs.10882         HMG-box containing protein muscle           5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptid muscle           5330         GGGGAGGGGC         3,00         0,45         Hs.89781         upstream binding transcrig RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein heterogeneous nuclear rit L           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear rit L           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         ilvB (bacterial acetolactate detactate detactat	: 1 (B5R.1)
DKFZp7610051 (from clone DKF	DKFZp7610051 (from closed containing protection   S328 ATGTTGCCCT   2,00   0,33 Hs.10882   HMG-box containing protection   S329 TGCTAAAAAAA   6,00   0,68 Hs.146550   myosin, heavy polypeptide muscle	
5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing protein 1           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptide 9, non-muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781         upstream binding transcription factor, RNA polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate synthase)-like           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily 1, group I, member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649         cytochrome c oxidase subunit VIc           5337 GGTGCGGCTGG         2,00         0,34 Hs.188882         Homo sapiens clone 23872 mRNA sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.170980         ESTs, Weakly similar to CBF1 interacting corepressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318         HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein	5328 ATGTTGCCCCT         2,00         0,33 Hs.10882         HMG-box containing prote           5329 TGCTAAAAAAA         6,00         0,68 Hs.146550 myosin, heavy polypeptid muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781 upstream binding transcription RNA polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804 DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730 heterogeneous nuclear rith           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880 ilvB (bacterial acetolactate           5334 CCAACCGTGCT         8,00         0,81 Hs.75207 glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623 nuclear receptor subfamily member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649 cytochrome c oxidase subsequence           5338 TTGTGATTAAT         2,00         0,34 Hs.188882 Homo sapiens clone 2387 sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.170980 ESTs, Weakly similar to Correpressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318 HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.11538 actin related protein 2/3 cut in related 2/3 cut in r	NA
5329 TGCTAAAAAAA         6,00         0,68 Hs.146550         myosin, heavy polypeptide 9, non-muscle           5330 GGGGAGGGGC         3,00         0,45 Hs.89781         upstream binding transcription factor, RNA polymeras           5331 GCCAGACACCC         3,00         0,45 Hs.3804         DKFZP564C1940 protein           5332 TACTCTTGGCA         14,00         1,11 Hs.2730         heterogeneous nuclear ribonucleoprotein L           5333 TAGGCCCAAGT         4,00         0,54 Hs.78880         ilvB (bacterial acetolactate synthase)-like           5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily 1, group I, member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649         cytochrome c oxidase subunit VIc           5337 GGTGCGGCTGG         2,00         0,34 Hs.188882         Homo sapiens clone 23872 mRNA sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.170980         ESTs, Weakly similar to CBF1 interacting corepressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318         HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.15344         ATRAP protein	5329         TGCTAAAAAAA         6,00         0,68         Hs.146550         myosin, heavy polypeptid muscle           5330         GGGGAGGGGC         3,00         0,45         Hs.89781         upstream binding transcrig RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear rit           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         ilvB (bacterial acetolactate           5334         CCAACCGTGCT         8,00         0,81         Hs.75207         glyoxalase I           5335         GGTGACCACCA         4,00         0,54         Hs.83623         nuclear receptor subfamily member 3           5336         GCATTTGACAG         2,00         0,34         Hs.74649         cytochrome c oxidase subsequence           5338         TTGTGATTAAT         2,00         0,34         Hs.18442         E-1 enzyme           5339         GGAGAGACAGG         2,00         0,34         Hs.170980         ESTs, Weakly similar to Correpressor           5340         TCAGTTTGTCA         10,00         0,92         Hs.15318         HS1 binding protein	
muscle   muscle   muscle   upstream binding transcription factor, RNA polymeras   DKFZP564C1940 protein   heterogeneous nuclear ribonucleoprotein   L   S331 TAGGCCAAGT   4,00   0,54 Hs.78880   itvB (bacterial acetolactate synthase)-like   S332 TAGGCCAAGT   4,00   0,54 Hs.78880   itvB (bacterial acetolactate synthase)-like   S334 CCAACCGTGCT   8,00   0,81 Hs.75207   glyoxalase   s335 GGTGACCACCA   4,00   0,54 Hs.83623   nuclear receptor subfamily 1, group I, member 3   member 3   s336 GCATTTGACAG   2,00   0,34 Hs.74649   cytochrome c oxidase subunit VIc   s337 GGTGCGGCTGG   2,00   0,34 Hs.188882   Homo sapiens clone 23872 mRNA   sequence   s338 TTGTGATTAAT   2,00   0,34 Hs.18442   E-1 enzyme   s339 GGAGAGACAGG   2,00   0,34 Hs.18442   E-1 enzyme   s340 TCAGTTTGTCA   10,00   0,92 Hs.15318   HS1 binding protein   s341 GTGCCTAGGGA   3,00   0,45 Hs.12854   ATRAP protein   s342 AGTATCTGGGA   3,00   0,45 Hs.11538   actin related protein 2/3 complex, subunit 1A (41 kD   s343 TCAATCAAGAT   9,00   0,87 Hs.75544   tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	S330   GGGAGGGGC   3,00   0,45   Hs.89781   Upstream binding transcription	ein 1
5330 GGGAGGGGC 3,00 0,45 Hs.89781 upstream binding transcription factor, RNA polymeras  5331 GCCAGACACCC 3,00 0,45 Hs.3804 DKFZP564C1940 protein  5332 TACTCTTGGCA 14,00 1,11 Hs.2730 heterogeneous nuclear ribonucleoprotein L  5333 TAGGCCCAAGT 4,00 0,54 Hs.78880 iiVB (bacterial acetolactate synthase)-like 5334 CCAACCGTGCT 8,00 0,81 Hs.75207 glyoxalase I  5335 GGTGACCACCA 4,00 0,54 Hs.83623 nuclear receptor subfamily 1, group I, member 3  5336 GCATTTGACAG 2,00 0,34 Hs.74649 cytochrome c oxidase subunit VIc 5337 GGTGCGGCTGG 2,00 0,34 Hs.188882 Homo sapiens clone 23872 mRNA sequence  5338 TTGTGATTAAT 2,00 0,34 Hs.18442 E-1 enzyme  5339 GGAGAGACAGG 2,00 0,34 Hs.170980 ESTs, Weakly similar to CBF1 interacting corepressor  5340 TCAGTTTGTCA 10,00 0,92 Hs.15318 HS1 binding protein 5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein actin related protein 2/3 complex, subunit 1A (41 kD)  5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5330         GGGGAGGGGC         3,00         0,45         Hs.89781         upstream binding transcription RNA polymeras           5331         GCCAGACACCC         3,00         0,45         Hs.3804         DKFZP564C1940 protein           5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear rik           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         ilvB (bacterial acetolactate           5334         CCAACCGTGCT         8,00         0,81         Hs.75207         glyoxalase I           5335         GGTGACCACCA         4,00         0,54         Hs.83623         nuclear receptor subfamily member 3           5336         GCATTTGACAG         2,00         0,34         Hs.74649         cytochrome c oxidase subsequence           5338         TTGTGATTAAT         2,00         0,34         Hs.18882         Homo sapiens clone 2387 sequence           5338         TTGTGATTAAT         2,00         0,34         Hs.170980         ESTs, Weakly similar to 0 corepressor           5340         TCAGTTTGTCA         10,00         0,92         Hs.15318         HS1 binding protein           5341         GTGCCTAGGGA         3,00         0,45         Hs.11538         ATRAP protein	e 9, non-
RNA polymeras   S331 GCCAGACACCC   3,00   0,45 Hs.3804   DKFZP564C1940 protein	RNA polymeras	
RNA polymeras   S331   GCCAGACACCC   3,00   0,45   Hs.3804   DKFZP564C1940 protein	RNA polymeras   S331 GCCAGACACCC   3,00   0,45 Hs.3804   DKFZP564C1940 protein   5332 TACTCTTGGCA   14,00   1,11 Hs.2730   heterogeneous nuclear rik   L   5333 TAGGCCCAAGT   4,00   0,54 Hs.78880   ilvB (bacterial acetolactate   5334 CCAACCGTGCT   8,00   0,81 Hs.75207   glyoxalase   1   5335 GGTGACCACCA   4,00   0,54 Hs.83623   nuclear receptor subfamily   member 3     1   1   1   1   1   1   1   1   1	ption factor,
TACTCTTGGCA 14,00 1,11 Hs.2730 heterogeneous nuclear ribonucleoprotein L  5333 TAGGCCCAAGT 4,00 0,54 Hs.78880 ilvB (bacterial acetolactate synthase)-like 5334 CCAACCGTGCT 8,00 0,81 Hs.75207 glyoxalase I  5335 GGTGACCACCA 4,00 0,54 Hs.83623 nuclear receptor subfamily 1, group I, member 3  5336 GCATTTGACAG 2,00 0,34 Hs.74649 cytochrome c oxidase subunit VIc 5337 GGTGCGGCTGG 2,00 0,34 Hs.188882 Homo sapiens clone 23872 mRNA sequence 5338 TTGTGATTAAT 2,00 0,34 Hs.18442 E-1 enzyme 5339 GGAGAGACAGG 2,00 0,34 Hs.170980 ESTs, Weakly similar to CBF1 interacting corepressor 5340 TCAGTTTGTCA 10,00 0,92 Hs.15318 HS1 binding protein 5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein actin related protein 2/3 complex, subunit 1A (41 kD 5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5332         TACTCTTGGCA         14,00         1,11         Hs.2730         heterogeneous nuclear rik L           5333         TAGGCCCAAGT         4,00         0,54         Hs.78880         ilvB (bacterial acetolactate ilvB (	
L	S333 TAGGCCCAAGT	
L	S333 TAGGCCCAAGT	bonucleoprotein
5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily 1, group I, member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649         cytochrome c oxidase subunit VIc           5337 GGTGCGGCTGG         2,00         0,34 Hs.18882         Homo sapiens clone 23872 mRNA sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.18442         E-1 enzyme           5339 GGAGAGACAGG         2,00         0,34 Hs.170980         ESTs, Weakly similar to CBF1 interacting corepressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318         HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.11538         actin related protein 2/3 complex, subunit 1A (41 kD           5343 TCAATCAAGAT         9,00         0,87 Hs.75544         tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5334 CCAACCGTGCT         8,00         0,81 Hs.75207         glyoxalase I           5335 GGTGACCACCA         4,00         0,54 Hs.83623         nuclear receptor subfamily member 3           5336 GCATTTGACAG         2,00         0,34 Hs.74649         cytochrome c oxidase substantial member 3           5337 GGTGCGGCTGG         2,00         0,34 Hs.188882         Homo sapiens clone 2387 sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.18442         E-1 enzyme           5339 GGAGAGACAGG         2,00         0,34 Hs.170980         ESTs, Weakly similar to 0 corepressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318         HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.11538         actin related protein 2/3 contact and 1A (41 kD)	·
5335 GGTGACCACCA 4,00 0,54 Hs.83623 nuclear receptor subfamily 1, group I, member 3 5336 GCATTTGACAG 2,00 0,34 Hs.74649 cytochrome c oxidase subunit VIc 5337 GGTGCGGCTGG 2,00 0,34 Hs.18882 Homo sapiens clone 23872 mRNA sequence 5338 TTGTGATTAAT 2,00 0,34 Hs.18442 E-1 enzyme 5339 GGAGAGACAGG 2,00 0,34 Hs.170980 ESTs, Weakly similar to CBF1 interacting corepressor 5340 TCAGTTTGTCA 10,00 0,92 Hs.15318 HS1 binding protein 5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein 5342 AGTATCTGGGA 3,00 0,45 Hs.11538 actin related protein 2/3 complex, subunit 1A (41 kD) 5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5335         GGTGACCACCA         4,00         0,54         Hs.83623         nuclear receptor subfamily member 3           5336         GCATTTGACAG         2,00         0,34         Hs.74649         cytochrome c oxidase sub cytochrom	e synthase)-like
member 3	member 3	
5336 GCATTTGACAG2,000,34 Hs.74649cytochrome c oxidase subunit VIc5337 GGTGCGGCTGG2,000,34 Hs.188882Homo sapiens clone 23872 mRNA sequence5338 TTGTGATTAAT2,000,34 Hs.18442E-1 enzyme5339 GGAGAGACAGG2,000,34 Hs.170980ESTs, Weakly similar to CBF1 interacting corepressor5340 TCAGTTTGTCA10,000,92 Hs.15318HS1 binding protein5341 GTGCCTAGGGA3,000,45 Hs.12854ATRAP protein5342 AGTATCTGGGA3,000,45 Hs.11538actin related protein 2/3 complex, subunit 1A (41 kD5343 TCAATCAAGAT9,000,87 Hs.75544tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5336 GCATTTGACAG         2,00         0,34 Hs.74649         cytochrome c oxidase substance           5337 GGTGCGGCTGG         2,00         0,34 Hs.188882         Homo sapiens clone 2387 sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.18442         E-1 enzyme           5339 GGAGAGACAGG         2,00         0,34 Hs.170980         ESTs, Weakly similar to Conceptressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318         HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.11538         actin related protein 2/3 contact to the conceptration of the	y 1, group I,
5337 GGTGCGGCTGG 2,00 0,34 Hs.188882 Homo sapiens clone 23872 mRNA sequence 5338 TTGTGATTAAT 2,00 0,34 Hs.18442 E-1 enzyme 5339 GGAGAGACAGG 2,00 0,34 Hs.170980 ESTs, Weakly similar to CBF1 interacting corepressor 5340 TCAGTTTGTCA 10,00 0,92 Hs.15318 HS1 binding protein 5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein 5342 AGTATCTGGGA 3,00 0,45 Hs.11538 actin related protein 2/3 complex, subunit 1A (41 kD) 5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5337 GGTGCGGCTGG         2,00         0,34 Hs.188882 Homo sapiens clone 2387 sequence           5338 TTGTGATTAAT         2,00         0,34 Hs.18442 E-1 enzyme           5339 GGAGAGACAGG         2,00         0,34 Hs.170980 ESTs, Weakly similar to Corepressor           5340 TCAGTTTGTCA         10,00         0,92 Hs.15318 HS1 binding protein           5341 GTGCCTAGGGA         3,00         0,45 Hs.12854 ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.11538 actin related protein 2/3 contact the contact	
Sequence   5338 TTGTGATTAAT   2,00   0,34 Hs.18442   E-1 enzyme   5339 GGAGACAGG   2,00   0,34 Hs.170980   ESTs, Weakly similar to CBF1 interacting corepressor   5340 TCAGTTTGTCA   10,00   0,92 Hs.15318   HS1 binding protein   5341 GTGCCTAGGGA   3,00   0,45 Hs.12854   ATRAP protein   5342 AGTATCTGGGA   3,00   0,45 Hs.11538   actin related protein 2/3 complex, subunit 1A (41 kD   1A (41	Sequence   S338 TTGTGATTAAT   2,00   0,34 Hs.18442   E-1 enzyme   E5339 GGAGAGACAGG   2,00   0,34 Hs.170980   ESTs, Weakly similar to Corepressor   S340 TCAGTTTGTCA   10,00   0,92 Hs.15318   HS1 binding protein   E341 GTGCCTAGGGA   3,00   0,45 Hs.12854   ATRAP protein   ATRAP	ounit VIc
Sequence   5338 TTGTGATTAAT   2,00   0,34 Hs.18442   E-1 enzyme   5339 GGAGACAGG   2,00   0,34 Hs.170980   ESTs, Weakly similar to CBF1 interacting corepressor   5340 TCAGTTTGTCA   10,00   0,92 Hs.15318   HS1 binding protein   5341 GTGCCTAGGGA   3,00   0,45 Hs.12854   ATRAP protein   5342 AGTATCTGGGA   3,00   0,45 Hs.11538   actin related protein 2/3 complex, subunit 1A (41 kD   1A (41	Sequence   S338 TTGTGATTAAT   2,00   0,34 Hs.18442   E-1 enzyme   E5339 GGAGAGACAGG   2,00   0,34 Hs.170980   ESTs, Weakly similar to Corepressor   S340 TCAGTTTGTCA   10,00   0,92 Hs.15318   HS1 binding protein   E341 GTGCCTAGGGA   3,00   0,45 Hs.12854   ATRAP protein   ATRAP	
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5339 GGAGAGACAGG 2,00 0,34 Hs.170980 ESTs, Weakly similar to CBF1 interacting corepressor  5340 TCAGTTTGTCA 10,00 0,92 Hs.15318 HS1 binding protein  5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein  5342 AGTATCTGGGA 3,00 0,45 Hs.11538 actin related protein 2/3 complex, subunit 1A (41 kD)  5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5339         GGAGAGACAGG         2,00         0,34         Hs.170980         ESTs, Weakly similar to Conceptessor           5340         TCAGTTTGTCA         10,00         0,92         Hs.15318         HS1 binding protein           5341         GTGCCTAGGGA         3,00         0,45         Hs.12854         ATRAP protein           5342         AGTATCTGGGA         3,00         0,45         Hs.11538         actin related protein 2/3 contact 1A (41 kD)	
corepressor  5340 TCAGTTTGTCA 10,00 0,92 Hs.15318 HS1 binding protein  5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein  5342 AGTATCTGGGA 3,00 0,45 Hs.11538 actin related protein 2/3 complex, subunit 1A (41 kD  5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	Corepressor     5340 TCAGTTTGTCA   10,00   0,92 Hs.15318   HS1 binding protein     5341 GTGCCTAGGGA   3,00   0,45 Hs.12854   ATRAP protein     5342 AGTATCTGGGA   3,00   0,45 Hs.11538   actin related protein 2/3 c   1A (41 kD	DBF1 interacting
5341 GTGCCTAGGGA3,000,45 Hs.12854ATRAP protein5342 AGTATCTGGGA3,000,45 Hs.11538actin related protein 2/3 complex, subunit 1A (41 kD5343 TCAATCAAGAT9,000,87 Hs.75544tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.11538         actin related protein 2/3 creation 1A (41 kD)	
5341 GTGCCTAGGGA 3,00 0,45 Hs.12854 ATRAP protein 5342 AGTATCTGGGA 3,00 0,45 Hs.11538 actin related protein 2/3 complex, subunit 1A (41 kD 5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	5341 GTGCCTAGGGA         3,00         0,45 Hs.12854         ATRAP protein           5342 AGTATCTGGGA         3,00         0,45 Hs.11538         actin related protein 2/3 creation 1A (41 kD)	
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5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	1A (41 kD	omplex, subunit
5343 TCAATCAAGAT 9,00 0,87 Hs.75544 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase		
monooxygenase	5343 TCAATCAAGAT   9,00  0,87 Hs.75544   tyrosine 3-monooxygenas	se/tryptophan 5-
100441 TOGO TOGOMAN TOTOLO OLOSIOS. 1/2004 VESICIE-ASSOCIATED MEMBRANE DIOTEIN O	5344 TGGCTGGGAAA 6,00 0,69 Hs.172684 vesicle-associated memb	rane protein 8
(endobrevin)		
5345 TAAGAAGCCCC 1,00 0,17 Hs.94318 ESTs		
	5346 CAGGCACTGAA 1,00 0,17 Hs.91065 hypothetical protein DKF2	Zp761B2423

<u></u>			- 4-		
5347	GGGTTTTTCTG	1,00			actin related protein 2/3 complex, subunit 2 (34 kD)
5348	GAGAAGACACG	1,00	0,17	Hs.8037	ESTs
5349	TGCAGAAGTAG	1,00	0,17	Hs.77770	adaptor-related protein complex 3, mu 2 subunit
5350	TTGTCCGGGCT	1,00	0,17	Hs.75064	tubulin-specific chaperone c
	GATTTTAATGT	1,00	0,17	Hs.7370	phosphotidylinositol transfer protein, beta
5352	CCTCTTGTAAT	1,00	0,17	Hs.66170	HSKM-B protein
5353	CCCTGGCTGTA	1,00	0,17	Hs.58127	ESTs
	TGATGTGGAAT	1,00		Hs.5687	protein phosphatase 1B (formerly 2C), magnesium-depe
5355	AAATGACTATA	1,00	0,17	Hs.43071	ESTs, Weakly similar to AF151900_1 CGI-142 protein [
5356	GAGGTTAGATT	1,00	0,17	Hs.283712	hypothetical protein
5357	ATGCTGTCTGC	1,00			Homo sapiens clone HQ0692
5358	AGTTGAGTCCT	1,00			S-adenosylmethionine decarboxylase 1
	CCCCACTAAAC	1,00			Homo sapiens mRNA; cDNA DKFZp434A1010 (from clone DK
5360	TGCTTGTGGTT	1,00	0,17	Hs.176600	WD-repeat protein
	TACGGGGGCCA	1,00			metastasis-associated 1-like 1
	CCGTGAAAAA	1,00		Hs.159448	
5363	TAACAAACCTG	1,00		Hs.15422	
	TTTCTGTGAAC	1,00			CGI-108 protein
	CAAATATCTTG	1,00		Hs.114404	
	GATGTGAAAAG	1,00			Homo sapiens mRNA; cDNA DKFZp434G0972 (from clone DK
5367	GCCACGTGGAG	1,00	0.17	Hs.103665	
	CACAAACGGTA	105,00			ribosomal protein S27 (metallopanstimulin 1)
5369	CGGAGGTGGGA	3,00	0,45	Hs.2491	DiGeorge syndrome critical region gene
5370	ATGGTGGTGGC	3,00	0.45	Hs.238030	secretory carrier membrane protein 2
	GCATATTAAAA	3,00			RAD23 (S. cerevisiae) homolog B
	GGTGAGACCTG	17,00			prostatic binding protein
	ACGCCCTGCTC	2,00		Hs.898	dystrophia myotonica-protein kinase
	AGGACTTCTGA	2,00			ESTs, Weakly similar to SFR7_HUMAN SPLICING FACTOR,
5375	ATCCGTGCCCT	6,00	0,70	Hs.141011	calmodulin 3 (phosphorylase kinase, delta)
5376	AAAAGAAACTT	15,00	1,21	Hs.172182	poly(A)-binding protein, cytoplasmic 1
5377	GGTGGATGTGC	5,00			methyl-CpG binding domain protein 3
5378	GTAGGGGCCTC	1,00		Hs.82208	acyl-Coenzyme A dehydrogenase, very long chain
5379	TAATTTGAAAA	1,00	0,18	Hs.6523	similar to rat smooth muscle protein SM-20
5380	AGGCCACCTCA	1,00	0,18	Hs.6084	frequenin (Drosophila) homolog
	ACTCTTGTTGG	1,00		Hs.5378	spondin 1, (f-spondin) extracellular matrix protein
5382	TGGAAGCTTTC	1,00	0,18	Hs.5308	ESTs

5393	ATACATAATAA	1,00	0.10	Ho 4004	impoduous of EDDD2 2
	GGGCCGCTCAG			Hs.4994 Hs.37656	transducer of ERBB2, 2
	TACCAAGCCAG	1,00		Hs.31388	KIAA0602 protein
	AATGCGGGAAA	1,00 1,00			
5366	AATGCGGGAAA	1,00	0,10	I S. 209030	ESTs, Moderately similar to ALUC HUMAN !!!! ALU CLAS
5397	AGGCTTTATGG	1 00	0.19	Un 24205	Human hbc647 mRNA sequence
-	TGATGAGTGCT	1,00 1,00		Hs.21657	
	TAAAACCCTAT	1,00			myosin, light polypeptide kinase
$\rightarrow$	GCCACAGCCAG	1,00			KIAA0599 protein
	GGGCTCCAGGA	1,00			F-box only protein 21
	AGCGGAGTCTG	1,00			phosphoglycerate mutase 1 (brain)
	CTTATGGTCCC	1,00			retinol dehydrogenase homolog
	GAAGTCATTTT	1,00	0,10	Ha 164470	ESTs, Weakly similar to AF161483_1
3334	GAAGICATITI	1,00	0, 10	ПS. 10447 о	HSPC134 [H.sapien
5305	TGTTTGGTTTC	1,00	0.18	He 161554	hypothetical protein FLJ20159
	AGAGCTCACTA	1,00	0,18		ESTs
	CCTGTAAATCC	1,00			hypothetical protein FLJ11271
	AGCTGTTCTGC	3,00	0,10	He 240405	heterogeneous nuclear ribonucleoprotein
3390	AGCIGITCIGC		0,40	П <b>5.24949</b> 5	A1
5399	GAGGCCAGTGA	3,00		Hs.2280	ribophorin I
5400	TGTAATCAATA	11,00	1,02	Hs.249495	heterogeneous nuclear ribonucleoprotein
					A1
	GATATCAGTCT	2,00		Hs.66394	ring finger protein 4
	CTTCTGCTGGG	2,00		Hs.17144	short-chain dehydrogenase/reductase 1
5403	AACTGCTTCAA	8,00	0,85	Hs.11538	actin related protein 2/3 complex, subunit
5 40 4	10007017070				1A (41 kD
	ACGGTGATGTC	5,00			ESTs
	GCCTCCTCCCA	14,00			muscle specific gene
	GCCCCTGCCTC	3,00			hypothetical protein DKFZp547H084
5407	TGCGGAGGCCC	3,00	0,47	Hs.25723	Sjogren's syndrome/scleroderma autoantigen 1
5408	CTTATGGTTGA	4,00	0,57	Hs.14084	ring finger protein 7
5409	TAATACTTTTG	2,00	0,35	Hs.90527	HSPC128 protein
5410	TAAGCATTAAA	2,00	0,35	Hs.8180	syndecan binding protein (syntenin)
	ACAGTGTTAAA	2,00	0,35	Hs.74649	cytochrome c oxidase subunit VIc
5412	TTTGTGGGCAG	2,00	0,35	Hs.39619	ESTs, Weakly similar to RCN1_HUMAN RETICULOCALBIN 1
5413	TGGTCCCTCTC	2,00	0,35	Hs.36587	protein phosphatase 1, regulatory
					subunit 7
_	GTTTCTAATAA	2,00			microtubule-associated protein 4
	CAATTGTAAAT	2,00			thioredoxin-like, 32kD
	GCACTTTGAGG	2,00			phospholipid scramblase 3
	ATCAAGGGTGT	32,00			ribosomal protein L9
	AGGGCGCAGA	4,00			SH3-domain GRB2-like 1
	GGCTGATGTGG	10,00			glycyl-tRNA synthetase
_	GCAGCCATCCG	81,00			ribosomal protein L28
	CTTCCTGTGAT	7,00	0,80	Hs.2533	aldehyde dehydrogenase 9 (gamma-
	AAGGAAGCTGC				aminobutyraldehyde d

5423	<b>GTGTTCCTCCG</b>	1,00	0.18	Hs.91299	unknown gene
	ACATAATAAAG	1,00		Hs.90077	TG-interacting factor (TALE family
3724	ACA IA IA AC	1,00	0,10	118.90077	homeobox)
5425	ACGTCTCTATT	1,00	0.18	Hs.8551	PRP4/STK/WD splicing factor
	TTGATGCCCTA	1,00		Hs.7871	hypothetical protein FLJ10081
	GACACCAACTA	1,00		Hs.77500	ubiquitin specific protease 4 (proto-
3427	GACACCAACTA	1,00	0, 10	Ins.77500	oncogene)
5/29	TGCCCAGCAAA	1,00	0.19	Hs.76297	G protein-coupled receptor kinase 6
	ATACATACTGT	1,00		Hs.74313	ESTs
	GAGGCCAATGC			Hs.6151	
		1,00			KIAA0235 protein ESTs
	TCTGCAGGGGA ACTACTAAATA	1,00		Hs.59509	
3432	ACIACIAAAIA	1,00	υ, ιο	Hs.5437	Tax1 (human T-cell leukemia virus type
5433	GCTGGTTCCTG	1,00	0.10	Hs.34516	I) binding pr Homo sapiens mRNA; cDNA
15455	GCIGGIICCIG	1,00	0,10	IDS.345 10	DKFZp434E0211 (from clone DK
5/3/	TGCCATATAAG	1,00	0 19	Hs.32271	hypothetical protein FLJ10846
	ACCCAATTTGT				ESTs
	ACCAAATATTT	1,00			
		1,00	0,10	HS.250641	tropomyosin 4
	GGACAGAACCC	1,00			hypothetical protein FLJ10889
	GTGATTATGAT	1,00			hypothetical protein FLJ10147
	TCACAGTGCCT	3,00			filamin B, beta (actin-binding protein-278)
5440	GCTCCACTGGA	3,00	0,47	Hs.75709	mannose-6-phosphate receptor (cation
E444	CTCCCCTCTTC		0.00	11-070444	dependent)
	GTGCCCTGTTG	5,00			NCK-associated protein 1
	CCCGGGGCCTC	2,00			nucleophosmin/nucleoplasmin 3
5443	CACCTGTCCTT	2,00		Hs.54457	CD81 antigen (target of antiproliferative antibody 1
5444	CTGGAGGCACA	2,00	0,35	Hs.283976	Homo sapiens clone TCBA00888 mRNA sequence
5445	GAGAGGCAGA	2,00	0.35	Hs.26412	ESTs, Weakly similar to KIAA0544
		-,	,,,,,		protein [H.sapiens]
5446	AATCCGACTCT	2,00	0,35	Hs.211577	kinectin 1 (kinesin receptor)
5447	CTTTTCAGCAA	4,00	0,58		tyrosine 3-monooxygenase/tryptophan 5-
i i		·	·		monooxygenase
5448	GGGCGGGGGG	2,00	0,36	Hs.99890	polymerase (DNA directed), delta 1,
					catalytic subuni
5449	GAGCGCAGCGA	2,00	0,36	Hs.83727	cleavage and polyadenylation specific
					factor 1, 160k
5450	TCCTGCCCTCA	2,00	0,36	Hs.80598	transcription elongation factor A (SII), 2
5451	TGTGGGAACCA	2,00		Hs.7750	Novel human gene mapping to
					chomosome 1
5452	CCAGATTTTGG	2,00	0,36	Hs.4788	nicastrin
5453	ATATTGTCAAA	2,00		Hs.3903	Homo sapiens mRNA; cDNA
					DKFZp762L106 (from clone DKF
5454	CACTGCATATG	2,00	0,36	Hs.196177	phosphorylase kinase, gamma 2 (testis)
5455	GAGCATAATAA	2,00			Wiskott-Aldrich syndrome-like
5456	CGCCGCCGGCT	66,00			ribosomal protein L35
	GGGGTGCTGTG	5,00		Hs.166161	<del></del>
<b> </b> 343/		3,001	0,01	113.100101	MYTICHINIC I

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E450	TETOTACTETO	44.00	4.00	11- 444004	epsilon
	TTTCTAGTTTG	11,00			membrane nucleoside transporter
	GGCAACGTGGT	6,00			Huntingtin interacting protein K
	AAGACAGTGGC	102,00			ribosomal protein L37a
5462	GAGGĞGAGGAA	1,00	0,19	Hs.83634	host cell factor C1 (VP16-accessory protein)
	TAAATGTTGAT	1,00			Human clone 23721 mRNA sequence
	CCTGCCACCCC	1,00		Hs.6133	ESTs, Highly similar to JC5772 tissue- specific calpa
5465	ACACTCTCCCC	1,00	0,19	Hs.57222	nurim (nuclear envelope membrane protein)
5466	GGAGTAAGGGG	1,00	0,19	Hs.5163	ESTs
5467	ATCGCGACACT	1,00	0,19	Hs.4864	KIAA0892 protein
5468	GCGAAAAAAA	1,00	0,19	Hs.4746	Homo sapiens mRNA; cDNA DKFZp761M16121 (from clone D
5469	TGTGCTGAGAG	1,00	0,19	Hs.284136	PRO2047 protein
5470	CTTTTAGGCCT	1,00	0,19	Hs.23202	ESTs
5471	CCTGTACTCCC	1,00		Hs.229434	
5472	CACTCACACAA	1,00			ribophorin I
5473	TGTATTCAGCA	1,00			hypothetical protein of unknown function
5474	TGGGAAACCTG	1,00			eukaryotic translation initiation factor 4 gamma, 1
5475	ATGCTGCCAAA	1,00	0,19	Hs.210749	proline synthetase co-transcribed (bacterial homolog
5476	CCAGCAGCTTC	1,00	0,19	Hs.200317	KIAA1404 protein
5477	AGACAGTAATA	1,00	0,19		acyl-Coenzyme A dehydrogenase, C-2 to C-3 short chai
5478	TATTTCAGTGA	1,00	0,19	Hs.12373	adenylate cyclase 6
5479	TGGCTAGATTT	1,00			UMP-CMP kinase
5480	CACTCTGGAAT	1,00		Hs.110707	H326
5481	CTGCCCTAGTA	1,00			chromosome 22 open reading frame 3
5482	AAGCTGAGTGG	7,00			heterogeneous nuclear ribonucleoprotein
5483	TTGCTGGAGAA	4,00	0,59		RNA binding protein; AT-rich element binding factor
5484	GCAGCTAATTT	3,00	0,49		GK001 protein
5485	AGAGCAAGTAC	3,00			small acidic protein
5486	CTTGATTAAAC	2,00		Hs.284265	Homo sapiens pRGR1 mRNA, partial cds
5487	GCCCGCAAGCT	2,00	0,36		bromodomain-containing 4
	TGAAAGTAACA	2,00		Hs.256583	interleukin enhancer binding factor 3, 90kD
5489	ACATCATCGAT	57,00	3,51		ribosomal protein L12
5490	TTTACAAAGAG	5,00			carboxypeptidase E
	AAGCCCAGGCT	3,00			DKFZP564N1362 protein
	ATCCACATCGC	8,00		Hs.119503	
	ACTGGCTGCTG	5,00			cytochrome c oxidase subunit VIc
	TGTGTGTTTGT	5,00			H1 histone family, member 0
	TTCCCCTTCCT	2,00			signal recognition particle receptor
					<u> </u>

				T	I/I-dealing proto
5406	GGGGAAGGGCA	2.00	0.27	U- 05077	('docking prote
	CCCTGAATCCC	2,00		Hs.65377	ESTS
5497	CCCTGAATCCC	2,00	0,37	HS.184592	Human clone A9A2BRB5
5400	AATCAACAATA	2 00		115 44040	(CAC)n/(GTG)n repeat-containing
	AATGAACAATA	2,00		Hs.11342	ninjurin 1
	GTTTGGCAGTG	6,00			hypothetical protein
	GAGGCCTCAGC	3,00			hypothetical protein FLJ20419
	TTCTGGCTGCG	7,00	0,86	Hs.119251	ubiquinol-cytochrome c reductase core protein I
	TGGTACACGTA	11,00	1,15	Hs.279574	CGI-39 protein
5503	GAGGGCCGGTG	6,00	0,78	Hs.36727	hypothetical protein FLJ10903
5504	GAAAAAATAAA	1,00	0,19	Hs.94925	dihydroorotate dehydrogenase
5505	GTTGTAAATAA	1,00	0,19	Hs.92033	Homo sapiens cDNA FLJ10181 fis, clone HEMBA1004227,
5506	AGGTGCCTCGG	1,00	0,19	Hs.84285	ubiquitin-conjugating enzyme E2I (homologous to yeas
5507	стпсстттс	1,00	0,19	Hs.80658	uncoupling protein 2 (mitochondrial, proton carrier)
5508	ATTTTTGCCCT	1,00	0.19	Hs.79372	retinoid X receptor, beta
	TTCCACCAACC	1,00		Hs.75618	RAB11A, member RAS oncogene family
	TAGGAGATTTT	1,00		Hs.74597	stromal interaction molecule 1
	TGGAAGAATGG	1,00		Hs.6774	ESTs
	TTCATTATAGG	1,00		Hs.6315	acetylserotonin O-methyltransferase-like
	GGCCTGTGTGA	1,00		Hs.4973	hypothetical protein
	AACTAACATTT	1,00		Hs.3297	ribosomal protein S27a
	CTGTATTAAAA	1,00		Hs.28264	Homo sapiens mRNA; cDNA
	_	1,00	0,10	113.20207	DKFZp564L0822 (from clone DK
5516	TTGCTATGAAA	1,00	0,19	Hs.26549	ESTs
5517	TTACTCTTTCT	1,00		Hs.2533	aldehyde dehydrogenase 9 (gamma- aminobutyraldehyde d
5518	ACTTATGTTTA	1,00	0.19	Hs.234896	aeminin
	CTGTTATAGGA	1,00			YME1 (S.cerevisiae)-like 1
	ATGGCTCACAC	1,00			DiGeorge syndrome critical region gene
5521	TTTTCCCACCA	1,00	0,19	Hs.153937	activated p21cdc42Hs kinase
5522	TTTCTGAAAAA	1,00	0,19	Hs.109646	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 6
5523	GACCTCACTGT	1,00	0,19		ESTs, Weakly similar to 2119210A mucin [H.sapiens]
5524	GTTTTTTTAA	1,00	0,19	Hs.10114	ESTs, Weakly similar to unnamed protein product [H.s
5525	стетссттете	5,00	0,70	Hs.6101	Human DNA sequence from clone 511E16 on chromosome 6
5526	CCACCCGAAT	15,00	1.41	Hs.74637	testis enhanced gene transcript
	TATGACTTAAT	7,00			Homo sapiens mRNA, clone:PO2ST9
	GCCCCAGCGAG	3,00			ADP-ribosylation factor binding protein
					GGA1
	CTGACCGGTGC	2,00			hematopoietic PBX-interacting protein
<u>5530</u>	TGTGTGCCACT	2,00	0,37	Hs.72925	chromosome 11 open reading frame 13

C504	IA ATTOOON OTO	0.00	0.07	111- 04000	FOT W. II
5531	AATTGCCACTG	2,00	0,37	Hs.61389	
5500		2.00	0.07		protein product [H.s
5532	ACAAAAAAA	2,00	0,37	Hs.274387	Homo sapiens mRNA; cDNA
					DKFZp434A1520 (from clone DK
	TGAATGTCAAG	2,00		Hs.230767	
5534	TACGAAGTTCT	2,00	0,37	Hs.19105	translocase of inner mitochondrial
					membrane 17 (yeas
5535	CTGAGTCTCCC	9,00	1,03	Hs.77269	guanine nucleotide binding protein (G
					protein), alph
	GAGGCGATCAG	4,00		Hs.30783	hypothetical protein FLJ20850
5537	GCCCCCACTC	3,00	0,50	Hs.75074	mitogen-activated protein kinase-
					activated protein k
5538	CACTCGTGTGA	3,00	0,50	Hs.146409	wingless-type MMTV integration site
EE30	CCACAACATCA	2.00	0.50	11- 420445	family, member 4
	GGAGAAGATGA	3,00		Hs.132415	·
$\overline{}$	TTTGGTCTTTT	3,00			hypothetical protein FLJ20625
	GGGGGACGGCT	7,00		Hs.21346	hypothetical protein LOC58481
	TAGTCCCTCTT	2,00		Hs.84264	acidic protein rich in leucines
	TATGCGTTTGG	2,00		Hs.76611	ESTs
5544	ATTTTGTGTCA	2,00	0,38	Hs.75056	adaptor-related protein complex 3, delta 1 subunit
5545	GACTCTGGTGC	12,00	1 26	Hs.2953	ribosomal protein S15a
	GCAGGTCAGCC	3,00		Hs.78950	branched chain keto acid
5546	GCAGGTCAGCC	3,00	0,51	ПS. / 090U	
5547	CACACTTTTT	2.00	0.54	No 74640	dehydrogenase E1, alpha pol
	CAGACTTTTT TTCAGTTGCTT	3,00		Hs.74649	cytochrome c oxidase subunit VIc
5546	ITCAGIIGCII	3,00	0,51	Hs.26700	Homo sapiens cDNA FLJ10309 fis, clone NT2RM2000287
5549	TAGAAACCAGA	3,00	0,51	Hs.194662	calponin 3, acidic
5550	ACTCGCTCTGT	3,00			laminin, alpha 5
5551	CTGGGTTAATA	99,00	6,16		ribosomal protein S19
	CCGTCCAAGGG	57,00			ribosomal protein S16
$\overline{}$	GAGTGAAAGAC	2,00		Hs.5811	hypothetical protein FLJ20467
	GAGCTCCACAG	2,00		Hs.3407	protein kinase (cAMP-dependent,
}		_,,	-,		catalytic) inhibitor
5555	AAAGTGGAAAA	2,00	0.38	Hs.203613	ESTs, Weakly similar to ALUB_HUMAN
( )			-,		IIII ALU CLASS B
5556	AATGGGGGTTA	1,00	0,19	Hs.94308	Homo sapiens cDNA FLJ10447 fis, clone NT2RP1000851
5557	CTTTGCTTTTT	1,00	0.10	Hs.84264	
	GTACGAATGGC	1,00		Hs.82065	acidic protein rich in leucines
	GIACGAAIGGC	1,00	0, 19	⊓5.02003	interleukin 6 signal transducer (gp130, oncostatin M
5559	TGAAGGTGGAT	1,00	0,19	Hs.7840	calcineurin binding protein 1
	GAATGAGCAAC	1,00		Hs.6686	ESTs
	GCTGCAAAGGA	1,00		Hs.61628	calcium binding atopy-related
					autoantigen 1
5562	CTAGATTCCCT	1,00	0,19	Hs.46783	ESTs
5563	TCTGTAGGCTG	1,00		Hs.38694	ESTs
5564	CAGGGCTCGCG	1,00		Hs.29288	Homo sapiens mRNA; cDNA
					DKFZp434P174 (from clone DKF

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<u> </u>	CCTGGGTCCTG	1,00			Homo sapiens cDNA FLJ20358 fis, clone HEP16618
	GAGAGGAAACT	1,00			hypothetical protein
5567	TGGTTCTATAT	1,00	0,19	Hs.26213	Human DNA sequence from clone RP3-447F3 on chromosom
5568	TGCCTATAGCC	1,00	0.19	Hs.258445	
	CACCCTGTACA	1,00			solute carrier family 29 (nucleoside transporters),
5570	AAGGAGAATGG	1,00	0,19	Hs.22119	Homo sapiens cDNA FLJ20318 fis, clone HEP08704
5571	CACGACTGTTC	1,00	0,19	Hs.184779	Homo sapiens mRNA; cDNA DKFZp586B1922 (from clone DK
5572	ATTITCTITA	1,00	·		TIA1 cytotoxic granule-associated RNA-binding protei
5573	AATGGTTAGCC	1,00	0,19		hypothetical protein PRO2219
5574	GTGGCATCACC	6,00	0,82	Hs.14317	hypothetical protein Nop10p
5575	AGCACCTCCAG	87,00	5,62	Hs.75309	eukaryotic translation elongation factor 2
5576	CCAAAATTAGG	3,00	0,51	Hs.239737	C-terminal binding protein 1
5577	TTTGTGACTGT	6,00	0,82	Hs.239737	C-terminal binding protein 1
5578	GTGCACTGAGC	23,00	2,01	Hs.181244	major histocompatibility complex, class I,
5579	CTCAAAAAAAA	4,00	0,63	Hs.165998	DKFZP564M2423 protein
5580	TTGGGGAAACA	2,00			biliverdin reductase A
5581	AGGCTGTGTTC	2,00	0,38	Hs.79	aminoacylase 1
5582	GGGGCTGTGGC	2,00	0,38	Hs.331	general transcription factor IIIC, polypeptide 1 (al
5583	тстстстстст	2,00	0,38		WS basic-helix-loop-helix leucine zipper protein
5584	СТТСТСТСТСС	2,00	0,38		ESTs, Weakly similar to cDNA EST yk415c12.5 comes fr
5585	AACGTGCAGGG	6,00	0,83		argininosuccinate synthetase
5586	GATAGTTGTGG	5,00			thyroid hormone receptor interactor 7
5587	CAAGGATCTAC	5,00			DKFZP586G1722 protein
5588	ATCAAGTTCGA	5,00	0,74	Hs.108082	ESTs, Weakly similar to Ydr472wp [S.cerevisiae]
	CAGGGGAGTGG	3,00		Hs.79396	N-methylpurine-DNA glycosylase
	CTCCAATAAAA	3,00	0,53	Hs.278559	talin
	CCTTATATTTG	3,00			tetratricopeptide repeat domain 3
	TTACGATGAAT	2,00	0,39	Hs.6335	phosphatidylinositol-4-phosphate 5-kinase, type II,
5593	CCACACACCGT	2,00	0,39	Hs.4877	CGI-51 protein
5594	GACTAGTGCGT	2,00		Hs.181551	
5595	GCAGCTCAGAT	1,00			Homo sapiens cDNA FLJ11021 fis, clone PLACE1003704,
5596	ACTCCAGCTGA	1,00	0,20	Hs.7763	vesicle docking protein p115
	GTGTCGCATCT	1,00		Hs.7734	Homo sapiens cDNA FLJ20684 fis, clone KAIA3469
5598	TGGAAAAAAA	1,00	0,20	Hs.61255	fructose-1,6-bisphosphatase 2
5599	GTAATTTAAAC	1,00		Hs.57664	Homo sapiens mRNA full length insert

	<del></del>	- <del></del> r			-DNA class EUDO
		- 1 22		11 1001	cDNA clone EURO
	CAGTATCCCAG	1,00		Hs.4994	transducer of ERBB2, 2
5601	AATCGCTAATA	1,00	0,20	Hs.47986	Homo sapiens mRNA; cDNA
					DKFZp586H051 (from clone DKF
	CCAGTTTGTAT	1,00			ephrin-A4
	AATAAAATTAG	1,00			mitogen-activated protein kinase 6
	AGGAGCCTTAG	1,00			A kinase (PRKA) anchor protein 8
5605	AATATGGTACA	1,00	0,20	Hs.236774	high-mobility group (nonhistone
					chromosomal) protein
5606	AATTTACTTCC	1,00	0,20	Hs.1742	IQ motif containing GTPase activating
					protein 1
	CCGGTTGGCAA	1,00			ESTs
	TTTGGAAAAAA	1,00		Hs.12482	glyceronephosphate O-acyltransferase
	GCTCATTAAAG	1,00		Hs.112237	
	TCCACGCACCA	4,00			hypothetical protein similar to mouse Fbw5
5611	ACTTACCTGCT	19,00			cytochrome c oxidase subunit VIb
5612	TCTGCTTACAG	7,00			ribosomal protein L15
5613	ATGACTCAAGG	5,00	0,76	Hs.239752	nuclear receptor subfamily 2, group F,
					member 6
5614	GATGAACACTG	2,00			CGI-130 protein
5615	TCGCCCAGGCG	2,00			ESTs
5616	ACAAGAATTGT	3,00	0,54	Hs.80919	synaptophysin-like protein
5617	TACCCCACCCT	9,00	1,13	Hs.7647	MYC-associated zinc finger protein
					(purine-binding t
	AGCCTGCAGAA	6,00			hypothetical protein R33729_1
5619	GGATTCCAGTT	3,00	0,54	Hs.5321	ARP3 (actin-related protein 3, yeast)
					homolog
5620	GGAGTCTAACT	3,00	0,54	Hs.240170	ESTs, Moderately similar to alternatively
					spliced pr
5621	GGAGTGTGCGT	2,00	0,40	Hs.4944	ESTs, Weakly similar to AIF1_HUMAN
					ALLOGRAFT INFLAMM
5622	GAAATCCGCAC	2,00	0,40	Hs.279854	mannosidase, alpha, class 2B, member
5000	TOTOTA 4 0 4 0 0	0.00	0.40	11 000000	
	TCTGTAACACC	2,00			butyrate-induced transcript 1
	TTCTGGCACTG	2,00			DKFZP566C134 protein
5625	GGCTCAAAACT	1,00	0,20	Hs.90625	Human DNA sequence from clone
5000	0.0000000000000000000000000000000000000	4 00	0.00	U- 77575	475B7 on chromosome Xq
	GTGGCCCCGGC	1,00		Hs.77575	ESTs
	TCTCTGCCTCT	1,00		Hs.77365	hypothetical protein FLJ11000
	TTCTGTAGCCC	1,00		Hs.5541	ATPase, Ca++ transporting, ubiquitous
	AATGGAGACTT	1,00		Hs.2943	signal recognition particle 19kD
	GAGGGCCTTCA	1,00			NICE-5 protein
	GTGAGGCCCCG	1,00			Homo sapiens cDNA FLJ20149 fis, clone COL08213
	TACCCACAGAG	1,00		Hs.26209	ESTs
5633	TTACCGTCCCC	1,00		Hs.25601	chromodomain helicase DNA binding protein 3
5634	GGAAGATGATG	1,00	0,20	Hs.19762	ESTs, Weakly similar to unknown

					[D.melanogaster]
5635	GAGGGAAATGG	1,00	0.20	Hs.12862	radical fringe (Drosophila) homolog
	AAGCTGTGTCT	1,00			ubiquitin-conjugating enzyme E2D 2
					(homologous to ye
	CACAGGGCCAT	1,00			ESTs, Weakly similar to PSF_HUMAN PTB-ASSOCIATED SPL
5638	GTAGATGCAAG	1,00			transcription factor 3 (E2A immunoglobulin enhancer
5639	TCAGCGGAGAA	1,00	0,20	Hs.100469	myeloid/lymphoid or mixed-lineage leukemia (trithora
5640	GTGACTGCCAC	5,00	0,78	Hs.84183	diptheria toxin resistance protein required for diph
5641	GGCCTGCTGCT	10,00	1,24	Hs.9634	ESTs, Highly similar to C10 [H.sapiens]
5642	AAGGAAGCAAT	4,00	0,67	Hs.194703	adaptor-related protein complex 4, mu 1 subunit
5643	AAATGCCACAC	8,00			reticulon 4
5644	TGTGTTGAGAG	150,00	10,78	Hs.275221	hypothetical protein FLJ20061
5645	CTGCTGAGTGA	6,00			hypothetical protein
5646	CCTGCTCCCTG	4,00	0,67	Hs.184601	solute carrier family 7 (cationic amino acid transpo
5647	GTAGAAAAGAA	2,00	0,40	Hs.75056	adaptor-related protein complex 3, delta 1 subunit
5648	AGCCTGTTGCA	2,00	0,40	Hs.182885	ESTs, Weakly similar to M03F8.2 [C.elegans]
5649	TGGAGGTGGGG	2,00	0,40	Hs.182625	VAMP (vesicle-associated membrane protein)-associate
5650	TGGCCCTCCAG	2,00	0,40	Hs.181015	signal transducer and activator of transcription 6,
5651	CCGTTCTGGAT	2,00	0,40	Hs.173638	Homo sapiens partial TCF-4 gene for T-cell transcrip
5652	GTGTTGGGGGT	3,00	0,55	Hs.55016	ESTs
5653	TGATGTTTGAC	5,00	0,79	Hs.75416	DAZ associated protein 2
5654	TCACAAGCAAA	23,00			nascent-polypeptide-associated complex alpha polypep
5655	TCAGAGAATAA	2,00	0,41	Hs.99486	ESTs
	CATCCCGTGAC	2,00			leukotriene A4 hydrolase
5657	TTAAGACTTCA	2,00	0,41	Hs.80562	gelsolin (amyloidosis, Finnish type)
5658	ACTTGCGAATA	2,00			fibrogenic lymphokine
	TCTGGGGAAAT	1,00	0,21	Hs.87417	cathepsin L2
5660	GTGTACCGGAT	1,00		Hs.8517	pleckstrin homology, Sec7 and coiled/coil domains 2
5661	ACGCTCATCGT	1,00	0,21	Hs.81217	frizzled (Drosophila) homolog 2
	TTCCTTTTTAC	1,00		Hs.75682	autoantigen
	ATGCAGAGATT	1,00		Hs.7137	clones 23667 and 23775 zinc finger protein
5664	TCACAAACTTC	1,00	0,21	Hs.61828	amyloid beta precursor protein-binding protein 1, 59
5665	AGACTATATTT	1,00	0,21	Hs.56542	X-prolyl aminopeptidase (aminopeptidase P) 1, solubl

Seequence   Seria   Sequence   Seria   Sequence   Seria   Se		<u> </u>	,			
Se67   TITCTGTTTTA	5666	CCTGAAAAGCT	1,00	0,21	Hs.3964	Homo sapiens clone 24877 mRNA sequence
5668   TTCTTTTTGA	5667	TITCTGTTTTA	1,00	0,21	Hs.31189	ESTs, Weakly similar to ALU2_HUMAN
5669   TIGTTGGTCAA	5668	TITCTTTTTGA	1.00	0.21	Hs 286236	
Se70   GGAGCCAGCTG						
5671 CAGAAGTGTCA         1,00         0,21 Hs.278441 KIAA0015 gene product           5672 ATGGGGAAGA         1,00         0,21 Hs.24989 ESTs         ESTs, Weakly similar to AF161483 HSPC134 [H.sapien           5674 CAAAAGCTTAT         1,00         0,21 Hs.208912 Novel human gene mapping to chomosome 22           5676 AAAGTGGCTAC         1,00         0,21 Hs.183668 glucuronidase, beta           5677 GCTAGTGAAAT         1,00         0,21 Hs.18406 endothelin converting enzyme.1           5678 CTAAACTTTTT         1,00         0,21 Hs.180919 inhibitor of DNA binding 2, dominar negative helix.           5679 CCTGTAACCCT         1,00         0,21 Hs.151777 Human translation initiation factor e 2alpha mRNA,           5680 CCTCCAGCCC         1,00         0,21 Hs.101174 microtubule-associated protein tau eukaryotic translation elongation fac gamma           5681 TGGGCAAAGCC         58,00         4,85 Hs.2186 eukaryotic translation elongation fac gamma           5684 AAGTGGGTGCC         6,00         0,92 Hs.119475 cold inducible RNA-binding protein homolog)           5685 TGCCAGGATT         2,00         0,41 Hs.91448 MKP-1 like protein tyrosine phosphit homologs           5687 GCTTAACCTGG         7,00         1,03 Hs.77508 glutamate dehydrogenase 1           5688 TGTCCCATT         7,00         1,03 Hs.77508 glutamate dehydrogenase 1           5689 GTGAACGAATA         6,00         0,92 Hs.8949 EsTs, Weakly simila	_					
5672   ATGGGGAGAGA   1,00   0,21   Hs.24989   ESTs   ESTs, Weakly similar to AF161483   Hs.22897   Hs.21941   AD021 protein   AD022 protein   AD02						
5673         GCGTGACTTCT         1,00         0,21         Hs.22897         ESTs, Weakly similar to AF161483 HSPC134 [H.saplen]           5674         CAAAAGCTTAT         1,00         0,21         Hs.21941         AD021 protein           5676         GCTTCGTGCTG         1,00         0,21         Hs.208912         Novel human gene mapping to chomosome 22           5676         AAAGTGGCTAC         1,00         0,21         Hs.183688 glucuronidase, beta           5677         GCTAGTGAAAT         1,00         0,21         Hs.181406 endothelin converting enzyme 1           5678         CTAAACTTTT         1,00         0,21         Hs.180919 inhibitor of DNA binding 2, dominan negative helix-           5679         CCTGTAACCT         1,00         0,21         Hs.151777         Human translation initiation factor eaghive helix-           5680         CCTCCAGCCC         1,00         0,21         Hs.146428 collagen, type V, alpha 1           5681         CTGTCTTTCT         1,00         0,21         Hs.101174         microtubule-associated protein tau           5682         TGGGCAAAGCC         58,00         4,85         Hs.2186         eukaryotic translation elongation factor agamma           5683         TGCGGCTGGTT         3,00         0,56         Hs.74617         hyrotin protein factor agam						
HSPC134   H.sapien						·
5674 CAAAAGCTTAT         1,00         0,21 Hs.21941         AD021 protein           5675 GCTTCGTGCTG         1,00         0,21 Hs.208912         Novel human gene mapping to chomosome 22           5676 AAAGTGGCTAC         1,00         0,21 Hs.183868 glucuronidase, beta           5677 GCTAGTGAAAT         1,00         0,21 Hs.181406 endothelin converting enzyme 1           5678 CTAAACTTTT         1,00         0,21 Hs.180919 inhibitor of DNA binding 2, dominan negative helix-           5679 CCTGTAACCCT         1,00         0,21 Hs.151777 Human translation initiation factor e2alpha mRNA,           5680 CCTCCAGCCCC         1,00         0,21 Hs.146428 collagen, type V, alpha 1           5681 CTGTCTGTTCT         1,00         0,21 Hs.101174 microtubule-associated protein tau eukaryotic translation elongation fac gamma           5682 TGGGCAAAGCC         58,00         4,85 Hs.2186 eukaryotic translation elongation fac gamma           5683 TGCGGCTGGTT         3,00         0,56 Hs.74617 homolog)           5684 AAGTGGGTGCC         6,00         0,92 Hs.119475 cold inducible RNA-binding protein           5685 TGCCCAGGATT         2,00         0,41 Hs.724 thyroid homone receptor, alpha (averythroblasti           5687 GCTTAACCTGG         7,00         1,03 Hs.77508 glutamate dehydrogenase 1           5688 GTATTCCCCTT         7,00         1,03 Hs.17176 poly(A)-binding protein, nuclear 1			',55	0,2.	110.22007	
5675         GCTTCGTGCTG         1,00         0,21         Hs.208912         Novel human gene mapping to chomosome 22           5676         AAAGTGGCTAC         1,00         0,21         Hs.183868         glucuronidase, beta           5677         GCTAGTGAAAT         1,00         0,21         Hs.181406         endothelin converting enzyme.1           5678         CTAAACTTTT         1,00         0,21         Hs.180919         inhibitor of DNA binding 2, dominan negative helix.           5679         CCTGTAACCCT         1,00         0,21         Hs.151777         Human translation initiation factor e 2alpha mRNA,           5680         CCTCCAGCCC         1,00         0,21         Hs.101174         microtubule-associated protein tau           5681         CTGTCTGTTCT         1,00         0,21         Hs.101174         microtubule-associated protein tau           5682         TGGGCAAAGCC         58,00         4,85         Hs.2186         eukaryotic translation elongation factor e 2alpha mRNA,           5683         TGCGGCTGGTT         3,00         0,56         Hs.74617         microtubule-associated protein tau           5683         TGCGGCTGGTT         3,00         0,56         Hs.74617         microtubule-associated protein tau           5684         AAGTGGGTGCC         6,00 </td <td>5674</td> <td>CAAAAGCTTAT</td> <td>1.00</td> <td>0.21</td> <td>Hs.21941</td> <td></td>	5674	CAAAAGCTTAT	1.00	0.21	Hs.21941	
Chomosome 22						
5676 AAAGTGGCTAC         1,00         0,21 Hs.183868 glucuronidase, beta           5677 GCTAGTGAAAT         1,00         0,21 Hs.181406 endothelin converting enzyme.1           5678 CTAAACTTTTT         1,00         0,21 Hs.180919 inhibitor of DNA binding 2, dominan negative helix-negative helix-n			.,	-,		
5677 GCTAGTGAAAT         1,00         0,21 Hs.181406 endothelin converting enzyme.1           5678 CTAAACTTTTT         1,00         0,21 Hs.180919 inhibitor of DNA binding 2, dominan negative helix-           5679 CCTGTAACCCT         1,00         0,21 Hs.151777 Human translation initiation factor ealpha mRNA,           5680 CCTCCAGCCC         1,00         0,21 Hs.146428 collagen, type V, alpha 1           5681 CTGTCTGTTCT         1,00         0,21 Hs.101174 microtubule-associated protein tau eukaryotic translation elongation factor in the elements of the elem	5676	AAAGTGGCTAC	1,00	0,21	Hs.183868	
5678 CTAAACTTTTT  1,00 0,21 Hs.180919 inhibitor of DNA binding 2, dominar negative helix- 5679 CCTGTAACCCT 1,00 0,21 Hs.151777 Human translation initiation factor e 2alpha mRNA, 5680 CCTCCAGCCCC 1,00 0,21 Hs.146428 collagen, type V, alpha 1 5681 CTGTCTGTTCT 1,00 0,21 Hs.101174 microtubule-associated protein tau 5682 TGGGCAAAGCC 58,00 4,85 Hs.2186 eukaryotic translation elongation factor of 2alpha mRNA, 5683 TGCGGCTGGTT 3,00 0,56 Hs.74617 dynactin 1 (p150, Glued (Drosophilation homolog) 5684 AAGTGGGTGCC 6,00 0,92 Hs.119475 cold inducible RNA-binding protein 5685 TGCCCAGGATT 2,00 0,41 Hs.91448 MKP-1 like protein tyrosine phosphatic hyroid homone receptor, alpha (averythroblasti) 5687 GCTTAACCTGG 7,00 1,03 Hs.77508 glutamate dehydrogenase 1 5688 GTATTCCCCTT 7,00 1,03 Hs.17176 poly(A)-binding protein, nuclear 1 5689 CTGAGCAATA 6,00 0,93 Hs.848 FK506-binding protein 4 (59kD) 5690 GTGATCATTAA 1,00 0,21 Hs.8949 ESTs, Weakly similar to AF126780 retinal short-cha sorbitol dehydrogenase 5692 GATGGAGCCT 1,00 0,21 Hs.7019 signal-induced proliferation-associa gene 1 5695 ACCAGGTCCAC 1,00 0,21 Hs.286078 E2k 5696 CTGGCCGACTT 1,00 0,21 Hs.286078 E2k 5697 GGGAACGGAGG 1,00 0,21 Hs.286078 E2k 5698 AATGGATTATT 1,00 0,21 Hs.267831 Rho GTPase activating protein 5 5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-jacyl-carrier-pr	5677	GCTAGTGAAAT	1,00			
negative helix-				0,21	Hs.180919	inhibitor of DNA binding 2. dominant
5679   CCTGTAACCCT   1,00   0,21   Hs.151777   Human translation initiation factor e			'	·		
2alpha mRNA,   5680   CCTCCAGCCCC   1,00   0,21   Hs.146428   collagen, type V, alpha 1   5681   CTGTCTGTTCT   1,00   0,21   Hs.101174   microtubule-associated protein tau   5682   TGGGCAAAGCC   58,00   4,85   Hs.2186   eukaryotic translation elongation fact gamma   5683   TGCGGCTGGTT   3,00   0,56   Hs.74617   dynactin 1 (p150, Glued (Drosophila homolog)   5684   AAGTGGGTGCC   6,00   0,92   Hs.119475   cold inducible RNA-binding protein   5685   TGCCCAGGATT   2,00   0,41   Hs.91448   MKP-1   like protein tyrosine phospha   TCTGTATCCCC   2,00   0,41   Hs.724   thyroid homone receptor, alpha (averythroblasti   5687   GCTTAACCTGG   7,00   1,03   Hs.77508   glutamate dehydrogenase 1   5689   CTTGAGCAATA   6,00   0,93   Hs.848   FK506-binding protein, nuclear 1   5689   CTTGAGCAATA   1,00   0,21   Hs.8949   ESTs, Weakly similar to AF126780   retinal short-cha   5691   AAAATCTGCCA   1,00   0,21   Hs.878   sorbitol dehydrogenase   5692   GATGGAGCCT   1,00   0,21   Hs.7019   signal-induced proliferation-associa gene 1   5695   ACCAGGTCCAC   1,00   0,21   Hs.286078   E2k   5695   ACCAGGTCCAC   1,00   0,21   Hs.285445   solute carrier family 5 (sodium-dependent vitamin tr   5697   GGGAACGGAGG   1,00   0,21   Hs.267831   Rho GTPase activating protein 5   5698   AATGGATTATT   1,00   0,21   Hs.247452   FabG (beta-ketoacyl-[acyl-carrier-protein   1,00   1,	5679	CCTGTAACCCT	1,00	0,21	Hs.151777	
5681 CTGTCTGTTCT1,000,21 Hs.101174microtubule-associated protein tau5682 TGGGCAAAGCC58,004,85 Hs.2186eukaryotic translation elongation fac gamma5683 TGCGGCTGGTT3,000,56 Hs.74617dynactin 1 (p150, Glued (Drosophila homolog)5684 AAGTGGGTGCC6,000,92 Hs.119475cold inducible RNA-binding protein5685 TGCCCAGGATT2,000,41 Hs.91448MKP-1 like protein tyrosine phosphe thyroid hormone receptor, alpha (av erythroblasti5687 GCTTAACCTGG7,001,03 Hs.77508glutamate dehydrogenase 15688 GTATTCCCCTT7,001,03 Hs.117176 poly(A)-binding protein, nuclear 15689 CTTGAGCAATA6,000,93 Hs.848FK506-binding protein 4 (59kD)5690 GTGATCATTAA1,000,21 Hs.8949ESTs, Weakly similar to AF126780 retinal short-cha5691 AAAATCTGCCA1,000,21 Hs.878sorbitol dehydrogenase5692 GATGGAGCCCT1,000,21 Hs.74649cytochrome c oxidase subunit VIc5693 CTCAGGAAGCT1,000,21 Hs.286078EZk5695 ACCAGGTCCAC1,000,21 Hs.286078EZk5696 CTGGCCGACTT1,000,21 Hs.286078EZk5697 GGGAACGGAGG1,000,21 Hs.267831Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452FabG (beta-ketoacyl-[acyl-carrier-protein)						
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TGGGCAAAGCC 58,00 4,85 Hs.2186 eukaryotic translation elongation factorized from the sum of the sum	5681	CTGTCTGTTCT	1,00			
5683 TGCGGCTGGTT 3,00 0,56 Hs.74617 dynactin 1 (p150, Glued (Drosophila homolog) 5684 AAGTGGGTGCC 6,00 0,92 Hs.119475 cold inducible RNA-binding protein 5685 TGCCCAGGATT 2,00 0,41 Hs.91448 MKP-1 like protein tyrosine phosphis 5686 TCTGTATCCCC 2,00 0,41 Hs.724 thyroid homone receptor, alpha (averythroblasti) 5687 GCTTAACCTGG 7,00 1,03 Hs.77508 glutamate dehydrogenase 1 5688 GTATTCCCCTT 7,00 1,03 Hs.117176 poly(A)-binding protein, nuclear 1 5689 CTTGAGCAATA 6,00 0,93 Hs.848 FK506-binding protein 4 (59kD) 5690 GTGATCATTAA 1,00 0,21 Hs.8949 ESTs, Weakly similar to AF126780 retinal short-cha 5691 AAAATCTGCCA 1,00 0,21 Hs.878 sorbitol dehydrogenase 5692 GATGGAGCCCT 1,00 0,21 Hs.74649 cytochrome c oxidase subunit VIc 5693 CTCAGGAAGCT 1,00 0,21 Hs.7019 signal-induced proliferation-associal gene 1 5694 GCTTTGTATCC 1,00 0,21 Hs.286078 E2k 5695 ACCAGGTCCAC 1,00 0,21 Hs.285445 solute carrier family 5 (sodium-dependent vitamin tr 5696 CTGGCCGACTT 1,00 0,21 Hs.274149 proline and glutamic acid rich nuclei protein 5697 GGGAACGGAGG 1,00 0,21 Hs.267831 Rho GTPase activating protein 5 5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr	5682	TGGGCAAAGCC	58,00			eukaryotic translation elongation factor 1
homolog    5684   AAGTGGGTGCC   6,00   0,92   Hs.119475   cold inducible RNA-binding protein     5685   TGCCCAGGATT   2,00   0,41   Hs.91448   MKP-1 like protein tyrosine phospha     5686   TCTGTATCCCC   2,00   0,41   Hs.724   thyroid hormone receptor, alpha (averythroblasti     5687   GCTTAACCTGG   7,00   1,03   Hs.77508   glutamate dehydrogenase 1     5688   GTATTCCCCTT   7,00   1,03   Hs.117176   poly(A)-binding protein, nuclear 1     5689   CTTGAGCAATA   6,00   0,93   Hs.848   FK506-binding protein 4 (59kD)     5690   GTGATCATTAA   1,00   0,21   Hs.8949   ESTs, Weakly similar to AF126780   retinal short-cha     5691   AAAATCTGCCA   1,00   0,21   Hs.878   sorbitol dehydrogenase     5692   GATGGAGCCCT   1,00   0,21   Hs.74649   cytochrome c oxidase subunit VIc     5693   CTCAGGAAGCT   1,00   0,21   Hs.7019   signal-induced proliferation-associa   gene 1     5694   GCTTTGTATCC   1,00   0,21   Hs.286078   E2k     5695   ACCAGGTCCAC   1,00   0,21   Hs.285445   solute carrier family 5 (sodium-dependent vitamin tr     5696   CTGGCCGACTT   1,00   0,21   Hs.274149   proline and glutamic acid rich nucleing     5697   GGGAACGGAGG   1,00   0,21   Hs.267831   Rho GTPase activating protein 5     5698   AATGGATTATT   1,00   0,21   Hs.247452   FabG (beta-ketoacyl-[acyl-carrier-protein     5697   GGGAACGGAGG   1,00   0,21   Hs.247452   FabG (beta-ketoacyl-[acyl-carrier-protein     5698   AATGGATTATT   1,00   0,21   Hs.247452   FabG (beta-ketoacyl-[acyl-carrier-protein     5699   5690						
5685 TGCCCAGGATT2,000,41 Hs.91448MKP-1 like protein tyrosine phospha5686 TCTGTATCCCC2,000,41 Hs.724thyroid hormone receptor, alpha (averythroblasti5687 GCTTAACCTGG7,001,03 Hs.77508glutamate dehydrogenase 15688 GTATTCCCCTT7,001,03 Hs.117176poly(A)-binding protein, nuclear 15689 CTTGAGCAATA6,000,93 Hs.848FK506-binding protein 4 (59kD)5690 GTGATCATTAA1,000,21 Hs.8949ESTs, Weakly similar to AF126780 retinal short-cha5691 AAAATCTGCCA1,000,21 Hs.74649cytochrome c oxidase subunit VIc5692 GATGGAGCCCT1,000,21 Hs.7019signal-induced proliferation-associa gene 15694 GCTTTGTATCC1,000,21 Hs.286078E2k5695 ACCAGGTCCAC1,000,21 Hs.285445solute carrier family 5 (sodium-dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452FabG (beta-ketoacyl-[acyl-carrier-protein)	5683	TGCGGCTGGTT	3,00			dynactin 1 (p150, Glued (Drosophila) homolog)
5685 TGCCCAGGATT2,000,41 Hs.91448MKP-1 like protein tyrosine phospha5686 TCTGTATCCCC2,000,41 Hs.724thyroid hormone receptor, alpha (averythroblasti5687 GCTTAACCTGG7,001,03 Hs.77508glutamate dehydrogenase 15688 GTATTCCCCTT7,001,03 Hs.117176poly(A)-binding protein, nuclear 15689 CTTGAGCAATA6,000,93 Hs.848FK506-binding protein 4 (59kD)5690 GTGATCATTAA1,000,21 Hs.8949ESTs, Weakly similar to AF126780 retinal short-cha5691 AAAATCTGCCA1,000,21 Hs.74649cytochrome c oxidase subunit VIc5692 GATGGAGCCCT1,000,21 Hs.7019signal-induced proliferation-associa gene 15694 GCTTTGTATCC1,000,21 Hs.286078E2k5695 ACCAGGTCCAC1,000,21 Hs.285445solute carrier family 5 (sodium-dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452FabG (beta-ketoacyl-[acyl-carrier-protein)	5684	AAGTGGGTGCC	6,00	0,92	Hs.119475	cold inducible RNA-binding protein
5686 TCTGTATCCCC 2,00 0,41 Hs.724 thyroid hormone receptor, alpha (averythroblasti) 5687 GCTTAACCTGG 7,00 1,03 Hs.77508 glutamate dehydrogenase 1 5688 GTATTCCCCTT 7,00 1,03 Hs.117176 poly(A)-binding protein, nuclear 1 5689 CTTGAGCAATA 6,00 0,93 Hs.848 FK506-binding protein 4 (59kD) 5690 GTGATCATTAA 1,00 0,21 Hs.8949 ESTs, Weakly similar to AF126780 retinal short-cha 5691 AAAATCTGCCA 1,00 0,21 Hs.878 sorbitol dehydrogenase 5692 GATGGAGCCCT 1,00 0,21 Hs.74649 cytochrome c oxidase subunit VIc 5693 CTCAGGAAGCT 1,00 0,21 Hs.7019 signal-induced proliferation-associa gene 1 5694 GCTTTGTATCC 1,00 0,21 Hs.286078 E2k 5695 ACCAGGTCCAC 1,00 0,21 Hs.285445 solute carrier family 5 (sodium-dependent vitamin tr 5696 CTGGCCGACTT 1,00 0,21 Hs.274149 proline and glutamic acid rich nuclei protein 5697 GGGAACGGAGG 1,00 0,21 Hs.267831 Rho GTPase activating protein 5 5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-protein-street family 5 (sodium-dependent vitamin tr	5685	TGCCCAGGATT	2,00	0,41	Hs.91448	MKP-1 like protein tyrosine phosphatase
5687 GCTTAACCTGG7,001,03 Hs.77508glutamate dehydrogenase 15688 GTATTCCCCTT7,001,03 Hs.117176poly(A)-binding protein, nuclear 15689 CTTGAGCAATA6,000,93 Hs.848FK506-binding protein 4 (59kD)5690 GTGATCATTAA1,000,21 Hs.8949ESTs, Weakly similar to AF126780_retinal short-cha5691 AAAATCTGCCA1,000,21 Hs.878sorbitol dehydrogenase5692 GATGGAGCCCT1,000,21 Hs.74649cytochrome c oxidase subunit VIc5693 CTCAGGAAGCT1,000,21 Hs.7019signal-induced proliferation-associa gene 15694 GCTTTGTATCC1,000,21 Hs.286078 E2k5695 ACCAGGTCCAC1,000,21 Hs.285445solute carrier family 5 (sodium- dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149proline and glutamic acid rich nuclea protein5697 GGGAACGGAGG1,000,21 Hs.267831 Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr	5686	TCTGTATCCCC	2,00	0,41	Hs.724	thyroid hormone receptor, alpha (avian
5688 GTATTCCCCTT7,001,03 Hs.117176 poly(A)-binding protein, nuclear 15689 CTTGAGCAATA6,000,93 Hs.848FK506-binding protein 4 (59kD)5690 GTGATCATTAA1,000,21 Hs.8949ESTs, Weakly similar to AF126780 retinal short-cha5691 AAAATCTGCCA1,000,21 Hs.878sorbitol dehydrogenase5692 GATGGAGCCCT1,000,21 Hs.74649cytochrome c oxidase subunit VIc5693 CTCAGGAAGCT1,000,21 Hs.7019signal-induced proliferation-associal gene 15694 GCTTTGTATCC1,000,21 Hs.286078 E2k5695 ACCAGGTCCAC1,000,21 Hs.285445solute carrier family 5 (sodium-dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831 Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-protein]	5687	GCTTAACCTGG	7,00	1,03	Hs.77508	
5689 CTTGAGCAATA6,000,93 Hs.848FK506-binding protein 4 (59kD)5690 GTGATCATTAA1,000,21 Hs.8949ESTs, Weakly similar to AF126780 retinal short-cha5691 AAAATCTGCCA1,000,21 Hs.878sorbitol dehydrogenase5692 GATGGAGCCCT1,000,21 Hs.74649cytochrome c oxidase subunit VIc5693 CTCAGGAAGCT1,000,21 Hs.7019signal-induced proliferation-associa gene 15694 GCTTTGTATCC1,000,21 Hs.286078 E2k5695 ACCAGGTCCAC1,000,21 Hs.285445solute carrier family 5 (sodium- dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831 Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-protein]	5688	GTATTCCCCTT	7,00	1,03	Hs.117176	poly(A)-binding protein, nuclear 1
retinal short-cha  5691 AAAATCTGCCA 1,00 0,21 Hs.878 sorbitol dehydrogenase  5692 GATGGAGCCCT 1,00 0,21 Hs.74649 cytochrome c oxidase subunit VIc  5693 CTCAGGAAGCT 1,00 0,21 Hs.7019 signal-induced proliferation-associa gene 1  5694 GCTTTGTATCC 1,00 0,21 Hs.286078 E2k  5695 ACCAGGTCCAC 1,00 0,21 Hs.285445 solute carrier family 5 (sodium-dependent vitamin tr  5696 CTGGCCGACTT 1,00 0,21 Hs.274149 proline and glutamic acid rich nuclei protein  5697 GGGAACGGAGG 1,00 0,21 Hs.267831 Rho GTPase activating protein 5  5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-protein-street]			6,00	0,93	Hs.848	
5692 GATGGAGCCCT1,000,21 Hs.74649cytochrome c oxidase subunit VIc5693 CTCAGGAAGCT1,000,21 Hs.7019signal-induced proliferation-associa gene 15694 GCTTTGTATCC1,000,21 Hs.286078 E2k5695 ACCAGGTCCAC1,000,21 Hs.285445solute carrier family 5 (sodium- dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831 Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr			1,00	0,21	Hs.8949	ESTs, Weakly similar to AF126780_1 retinal short-cha
5692 GATGGAGCCCT1,000,21 Hs.74649cytochrome c oxidase subunit VIc5693 CTCAGGAAGCT1,000,21 Hs.7019signal-induced proliferation-associal gene 15694 GCTTTGTATCC1,000,21 Hs.286078 E2k5695 ACCAGGTCCAC1,000,21 Hs.285445 solute carrier family 5 (sodium-dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149 proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831 Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-protein]						sorbitol dehydrogenase
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5694 GCTTTGTATCC1,000,21 Hs.286078 E2k5695 ACCAGGTCCAC1,000,21 Hs.285445 solute carrier family 5 (sodium-dependent vitamin tr5696 CTGGCCGACTT1,000,21 Hs.274149 proline and glutamic acid rich nuclei protein5697 GGGAACGGAGG1,000,21 Hs.267831 Rho GTPase activating protein 55698 AATGGATTATT1,000,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-protein protein 5)	5693	CTCAGGAAGCT	1,00			signal-induced proliferation-associated
5695ACCAGGTCCAC1,000,21Hs.285445solute carrier family 5 (sodium-dependent vitamin tr5696CTGGCCGACTT1,000,21Hs.274149proline and glutamic acid rich nuclei protein5697GGGAACGGAGG1,000,21Hs.267831Rho GTPase activating protein 55698AATGGATTATT1,000,21Hs.247452FabG (beta-ketoacyl-[acyl-carrier-protein 5)			1,00	0,21	Hs.286078	
dependent vitamin tr  5696 CTGGCCGACTT 1,00 0,21 Hs.274149 proline and glutamic acid rich nuclei protein  5697 GGGAACGGAGG 1,00 0,21 Hs.267831 Rho GTPase activating protein 5  5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr	5695	ACCAGGTCCAC				
5696 CTGGCCGACTT 1,00 0,21 Hs.274149 proline and glutamic acid rich nuclei protein  5697 GGGAACGGAGG 1,00 0,21 Hs.267831 Rho GTPase activating protein 5  5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr						
protein  5697 GGGAACGGAGG 1,00 0,21 Hs.267831 Rho GTPase activating protein 5  5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr	5696	CTGGCCGACTT	1,00	0,21	Hs.274149	
5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr						protein
5698 AATGGATTATT 1,00 0,21 Hs.247452 FabG (beta-ketoacyl-[acyl-carrier-pr				0,21	Hs.267831	Rho GTPase activating protein 5
			1,00	0,21	Hs.247452	FabG (beta-ketoacyl-[acyl-carrier-protein] reductase
	5699	AAGATAAACTC	1,00	0,21	Hs.19015	ESTs, Weakly similar to NR54_HUMAN

5700	TGGCTGGCCAC	1,00	0,21	Hs.174142	colony stimulating factor 1 receptor, formerly McDon
5701	CTTGTGATCCC	1,00	0.21	Hs 173042	KIAA1143 protein
	GTTAAAAAAAA	1,00			centaurin-alpha
	TGGGGCCGCAG	1,00		Hs.126759	
	TCTTTTTGGGG	1,00		Hs.120854	
	CTITIGTTITC	1,00			thyroid hormone receptor-associated
					protein, 150 kDa
5/06	CATAGTTTTAG	1,00	0,21	Hs.106823	H.sapiens gene from PAC 426l6, similar to syntaxin 7
5707	GCTGATTGGCA	1,00	0,21	Hs.104476	ESTs
5708	TGGTTGCGACA	1,00	0,21	Hs.101408	branched chain aminotransferase 2, mitochondrial
5709	TTACTAAATGG	6,00	0,94	Hs.155560	calnexin
5710	TACTAATAAAA	5,00	0,83		ESTs, Weakly similar to F15D4.3 [C.elegans]
5711	CCTTCTGGTGG	3,00	0.58		protein tyrosine kinase 9
	ATGACACTCAC	4,00			destrin (actin depolymerizing factor)
	GCTCCCAGACT	7,00		Hs.5097	synaptogyrin 2
$\overline{}$	GGAAGGGAGGC	6,00			hypothetical protein FLJ20568
	TAACCCAGCAG	3,00	0.58	Hs.7043	succinate-CoA ligase, GDP-forming,
					alpha subunit
	TAAAAAAAAA	17,00	·	Hs.98658	budding uninhibited by benzimidazoles 1 (yeast homol
5717	TTTTGTAAATA	2,00	0,43	Hs.75607	myristoylated alanine-rich protein kinase C substrat
5718	GĢCĄTTĠTTCA	2,00	0,43	Hs.3128	polymerase (RNA) II (DNA directed) polypeptide H
5719	TTGGGTTTTGA	2,00	0,43		ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY
5720	CCTGGATAAAT	2,00	0,43		Human Chromosome 16 BAC clone CIT987SK-A-61E3
5721	GGCAATATGGT	2,00	0,43	Hs.160823	
5722	GAAAAGAGATT	1,00		Hs.92145	
5723	TCTGTTAATAA	1,00		Hs.89434	
5724	TACATTCACCT	1,00			D123 gene product
5725	GTTCTCCCTTC	1,00	0,22	Hs.49657	Homo sapiens mRNA; cDNA DKFZp547N084 (from clone DKF
5726	CTGTGCTCTAA	1,00	0.22	Hs.42500	ADP-ribosylation factor-like 5
	TGGGCTGGGGT	1,00		Hs.3416	adipose differentiation-related protein
	GAAGAACAAGT	1,00		Hs.28491	spermidine/spermine N1-
		','	-,		acetyltransferase
5729	GGAAGATGTTC	1,00	0,22	Hs.284158	Homo sapiens mRNA; cDNA
5720	TTAATAAAATA	4.00	0.00	Ua 2707	DKFZp762B195 (from clone DKF
		1,00		Hs.2707	G1 to S phase transition 1
	ACCTTGGGCAA	1,00		Hs.19554	chromosome 1 open reading frame 2
	TGTACCTAACT	1,00			ESTs
	TGGCTTATTAA	1,00		Hs.18021	hypothetical protein FLJ20446
0/34	TTGTGATACTA	1,00	<u>U,22</u>	Hs.175417	LSIS

5736 TGTTTAATACA  1,00  0,22 Hs.143601 Homo sapiens cDNA FLJ20678 fis, clon KANA4163  5737 CACCACCACAC  1,00  0,22 Hs.123319 EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SE  5738 GGGAGCACTAA  1,00  0,22 Hs.175802 CGI-43 protein  5739 GGGAGGAGGGT  1,00  0,22 Hs.1717502 CGI-43 protein  5739 GGGAGGAGGGT  1,00  0,22 Hs.1717502 CGI-43 protein  5739 GGGAGGAGGGT  1,00  0,22 Hs.1717501 ESTs, Moderately similar to AF151825_ CGI-67 protei  5740 TGGACCCCCG  3,00  0,59 Hs.173501 ESTs, Moderately similar to AF151825_ CGI-67 protei  5741 GTACTGTATGC  4,00  0,73 Hs.180446 karyopherin (importin) beta 1  1,70 Hs.76159  ATPase, H+ transporting, lysosomal (vacuolar proton myxovirus (influenza) resistance 1, homolog of murin  5744 AGTGCCGTGTG  2,00  0,43 Hs.146409 imposer proton myxovirus (influenza) resistance 1, homolog of murin  5745 GCCTGAGGGGC  2,00  0,43 Hs.146409 imposer, the full milly member 4  5746 TCTCAATTCTT  5,00  0,86 Hs.146409 imposer, the full milly member 4  5747 TTCATTATAAT  9,00  1,32 Hs.250655 prothymosin, alpha (gene sequence 28) from subunit, alpha type.  5749 ATGGAAAGGAA  2,00  0,44 Hs.243901 Homo sapiens cDNA FLJ20738 fis, clone HEP08257  5750 TGCTGCTTGAA  2,00  0,44 Hs.243901 Homo sapiens cDNA FLJ20738 fis, clone HEP08257  TTTTCTGAGTG  1,00  0,22 Hs.82338  POP4 (processing of precursor, S. cerevisiae) homol  5754 TGATACTACT  1,00  0,22 Hs.75650 TLPS-induced TNF-alpha factor  5755 TCAAGTCCAGA  1,00  0,22 Hs.75657  CAAGCCACAACTGTG  1,00  0,22 Hs.75657  CAGGCAGAACAGA  1,00  0,22 Hs.75675  Solute carrier family 20 (phosphate transporter), me  5759 GAAGAATTGGA  1,00  0,22 Hs.6179  DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k  5759 GAAGGACACAT  1,00  0,22 Hs.52768 Forton soring nexin 12  5769 GAAGGCACCAT  1,00  0,22 Hs.52767  CHARLES STA STANA TOR TOR SIMILAT TOR	5735	GGCTTTCCCTG	1,00	0.22	He 171802	ESTs, Weakly similar to hypothetical
KAIA4163	5735	GGCTTTCCCTG	1,00			protein [H.sapi
ALU SUBFAMILY J SE	5736	TGTTTAATACA				KAIA4163
5739   GGAGGAGGGT   1,00   0,22   Hs.112159   ESTs   TGGACCCCCG   3,00   0,59   Hs.173501   ESTs, Moderately similar to AF151825_CGI-67 protei   CGI-67						ALU SUBFAMILY J SE
5740         TGGACCCCCG         3,00         0,59         Hs.173501         ESTs, Moderately similar to AF151825_CGI-67 protei           5741         GTACTGTATGC         4,00         0,73         Hs.180446         karyopherin (importin) beta 1           5742         GTGTGTTTGTA         19,00         2,26         Hs.118787         transforming growth factor, beta-induced 68kD           5743         CGCAGTGTCCT         13,00         1,70         Hs.76159         ATPase, H+ transporting, lysosomal (vacuolar proton myxovirus (influenza) resistance 1, homolog of murin           5744         AGTGCCGTGTG         2,00         0,43         Hs.14927         ESTs, Moderately similar to TNRC_HUMAN LYMPHOTOXIN-B           5745         GCCTGAGGGGC         2,00         0,43         Hs.146409         wingless-type MMTV integration site family, member 4           5746         TCTCAATTCTT         5,00         0,86         Hs.146409         wingless-type MMTV integration site family, member 4           5748         GATGAGTCTG         10,00         1,32         Hs.233952         prothymosin, alpha (gene sequence 28)           5749         ATGGAAAGGAA         2,00         0,44         Hs.212152         APMCF1 protein           5750         TGGTTGGAT         1,00         0,22         Hs.82238         virumin A responsive; cytoskeleton rela	5738	GGGAGCATTAA		0,22	Hs.117582	CGI-43 protein
CGI-67 protein	5739	GGGAGGAGGGT	1,00			
5742         GTGTGTTTGTA         19,00         2,26         Hs.118787 transforming growth factor, beta-induced 68kD           5743         CGCAGTGTCCT         13,00         1,70         Hs.76159         ATPase, H+ transporting, lysosomal (vacuolar proton)           5744         AGTGCCGTGTG         2,00         0,43         Hs.76391         myxovirus (influenza) resistance 1, homolog of murin homolog of murin           5745         GCCTGAGGGGC         2,00         0,43         Hs.14927         ESTs, Moderately similar to TNRC_HUMAN LYMPHOTOXIN-B           5746         TCTCAATTCTT         5,00         0,86         Hs.146409 wingless-type MMTV integration site family, member 4           5747         TTCATTATAAT         9,00         1,32         Hs.250655 prothymosin, alpha (gene sequence 28) proteasome (prosome, macropain) subunit, alpha type,           5749         ATGGAAAGGAA         2,00         0,44         Hs.243901 homo sapiens cDNA FLJ20738 fis, clone HEP08257           5750         TGTTAGCCTGT         1,00         0,22         Hs.92384         PMCF1 protein           5752         TTTTCTGAGTG         1,00         0,22         Hs.82238         POP4 (processing of precursor, Scerevisiae) homol           5753         ACCAAACTGTG         1,00         0,22         Hs.77608         splicing factor, arginine/serine-rich 9 <t< td=""><td>5740</td><td>TGGACCCCCCG</td><td>3,00</td><td></td><td></td><td>CGI-67 protei</td></t<>	5740	TGGACCCCCCG	3,00			CGI-67 protei
S8kD	5741	GTACTGTATGC	4,00			
(vacuolar proton   F744   AGTGCCGTGTG   2,00   0,43   Hs.76391   myxovirus (influenza) resistance 1, homolog of murin   ESTs, Moderately similar to TNRC HUMAN LYMPHOTOXIN-B   F746   TCTCAATTCTT   5,00   0,86   Hs.146409   wingless-type MMTV integration site family, member 4   F747   TTCATTATAAT   9,00   1,32   Hs.250655   prothymosin, alpha (gene sequence 28)   F748   GATGAGTCTCG   10,00   1,43   Hs.233952   proteasome (prosome, macropain)   subunit, alpha type,   F749   ATGGAAAGGAA   2,00   0,44   Hs.243901   Homo sapiens cDNA FLJ20738 fis, clone   HEP08257   HEP08257   HEP08257   TGTTAGCCTGT   1,00   0,22   Hs.82238   POP4 (processing of precursor , S. cerevisiae)   homol   F752   TGTTAGCCTGT   1,00   0,22   Hs.82238   POP4 (processing of precursor , S. cerevisiae)   homol   F755   TGAATACTACT   1,00   0,22   Hs.76507   LPS-induced TNF-alpha factor   S755   TGAATCCAGA   1,00   0,22   Hs.75867   solute carrier family 20 (phosphate transporter), me   F756   TGTACTTCCTA   1,00   0,22   Hs.6179   DEAD/H (Asp-Glu-Ala-Asp/His)   box polypeptide 17 (72k   F759   GAGGCAGAAGA   1,00   0,22   Hs.55158   ESTs, Weakly similar to weak similarity to TPR domai   S760   GTCTTCTTAAT   1,00   0,22   Hs.26706   CGI-121 protein   GAAGGCACCTT   1,00   0,22   Hs.26708   DNAJ domain-containing   S761   GAAGGCACCTT   1,00   0,22   Hs.26708   CGI-121 protein   GAAGGCACCTC   1,00   0,22   Hs.26708   CGI-121 protein   GAAGGCACCTC   1,00   0,22   Hs.267037   Cromosome 22 open reading frame 2   GAGGCACCAGGGACC   1,00   0,22   Hs.205421   EST   GAGCGCCCAGGGACC   1,00   0,22   Hs.205421   EST   GAGCGCCCAGGGACC   1,00   0,22   Hs.205421   EST   GAGCGCCCAGGACCCACCACCACCACCACCACCACCACCA					•	68kD
homolog of murin   FS745   GCCTGAGGGGC   2,00   0,43   Hs.14927   ESTs, Moderately similar to   TNRC_HUMAN LYMPHOTOXIN-B   TNRC	5743	CGCAGTGTCCT				(vacuolar proton
5745         GCCTGAGGGGC         2,00         0,43         Hs.14927         ESTs, Moderately similar to TNRC_HUMAN LYMPHOTOXIN-B           5746         TCTCAATTCTT         5,00         0,86         Hs.146409         wingless-type MMTV integration site family, member 4           5747         TTCATTATAAT         9,00         1,32         Hs.250655         prothymosin, alpha (gene sequence 28)           5748         GATGAGTCTCG         10,00         1,43         Hs.233952         proteasome (prosome, macropain) subunit, alpha type,           5749         ATGGAAAGGAA         2,00         0,44         Hs.243901         Homo sapiens cDNA FLJ20738 fis, clone HEP08257           5750         TGCTGCTTGAA         2,00         0,44         Hs.243901         Homo sapiens cDNA FLJ20738 fis, clone HEP08257           5751         TGTTAGCCTGT         1,00         0,22         Hs.92384         vitamin A responsive; cytoskeleton related           5752         TTTTCTGAGTG         1,00         0,22         Hs.7660         PDP4 (processing of precursor , S. cerevisiae) homol           5753         ACCAAACTGTG         1,00         0,22         Hs.76507         LPS-induced TNF-alpha factor           5754         TGAGTCCAGA         1,00         0,22         Hs.75867         solute carrier family 20 (phosphate transporter), me	5744	AGTGCCGTGTG	2,00	0,43	Hs.76391	
TNRC_HUMAN LYMPHOTOXIN-B   TNRC_HUMAN LYMPHOTOXIN-B   TOTCAATTCTT   5,00   0,86   Hs. 146409   wingless-type MMTV integration site   family, member 4   family, member 4   5747   TTCATTATAAT   9,00   1,32   Hs. 250655   prothymosin, alpha (gene sequence 28)   proteasome (prosome, macropain)   subunit, alpha type,   5749   ATGGAAAGGAA   2,00   0,44   Hs. 243901   Homo sapiens cDNA FLJ20738 fis, clone   HEP08257   TGCTGCTTGAA   2,00   0,44   Hs. 12152   APMCF1 protein   5751   TGTTAGCCTGT   1,00   0,22   Hs. 92384   vitamin A responsive; cytoskeleton related   5752   TTTTCTGAGTG   1,00   0,22   Hs. 7638   POP4 (processing of precursor , S. cerevisiae)   homol   5753   ACCAAACTGTG   1,00   0,22   Hs. 76507   LPS-induced TNF-alpha factor   5755   TCAAGTCCAGA   1,00   0,22   Hs. 75867   LPS-induced TNF-alpha factor   5756   TGTACTTCCTA   1,00   0,22   Hs. 75485   ornithine aminotransferase (gyrate atrophy)   5757   GAGGCAGAAGA   1,00   0,22   Hs. 6179   DEAD/H (Asp-Glu-Ala-Asp/His)   box   polypeptide 17 (72k   TST   TST   TCATAGTC   1,00   0,22   Hs. 55158   EST s, Weakly similar to weak similarity   to TPR domai   5769   GAAGGATTGGA   1,00   0,22   Hs. 279884   DNAJ domain-containing   5769   GATGAGCGGCT   1,00   0,22   Hs. 26023   KIAA1041 protein   5763   GATGAGCGGCT   1,00   0,22   Hs. 26023   KIAA1041 protein   5764   TGACTGCTGCT   1,00   0,22   Hs. 27637   chromosome 22 open reading frame 2   5764   TGACTGCTGCT   1,00   0,22   Hs. 205421   EST   CCCAGGGACC   1,00   0,22   Hs. 205421   EST   CCCAGGACCAGA   1,00   0,22   Hs. 205421   EST   CCCAGGACCAGA   1,00   0,22   Hs. 205421   EST   CCCAGGACCAGACCAGACCAGACAGACAGACAGACAGAC	5745	CCTGAGGGGC	2.00	0.43	He 14027	
family, member 4   family member 4			2,00			TNRC_HUMAN LYMPHOTOXIN-B
5748 GATGAGTCTCG         10,00         1,43 Hs.233952 proteasome (prosome, macropain) subunit, alpha type,           5749 ATGGAAAGGAA         2,00         0,44 Hs.243901 Homo sapiens cDNA FLJ20738 fis, clone HEP08257           5750 TGCTGCTTGAA         2,00         0,44 Hs.12152 APMCF1 protein           5751 TGTTAGCCTGT         1,00         0,22 Hs.92384 vitamin A responsive; cytoskeleton related           5752 TTTTCTGAGTG         1,00         0,22 Hs.82238 POP4 (processing of precursor , S. cerevisiae) homol           5753 ACCAAACTGTG         1,00         0,22 Hs.77608 splicing factor, arginine/serine-rich 9           5754 TGAATACTACT         1,00         0,22 Hs.76507 LPS-induced TNF-alpha factor           5755 TCAAGTCCAGA         1,00         0,22 Hs.75867 solute carrier family 20 (phosphate transporter), me           5756 TGTACTTCCTA         1,00         0,22 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k           5758 TGCCTATAGTC         1,00         0,22 Hs.55158 ESTs, Weakly similar to weak similarity to TPR domai           5759 GAAAGATTGGA         1,00         0,22 Hs.279884 DNAJ domain-containing           5760 GTCTTCTTAAT         1,00         0,22 Hs.26706 CGI-121 protein           5761 GAAGGCACCAT         1,00         0,22 Hs.26703 CHromosome 22 open reading frame 2           5763 GATGAGCGGCT         1,00         0,22 Hs.214372 ESTs           5765 GCCCAGGGACC<	5746	TCTCAATTCTT	5,00	0,86	Hs.146409	
Subunit, alpha type,	5747	TTCATTATAAT	9,00	1,32	Hs.250655	prothymosin, alpha (gene sequence 28)
HEP08257   TGCTGCTTGAA   2,00   0,44   Hs.12152   APMCF1 protein   vitamin A responsive; cytoskeleton related   POP4 (processing of precursor , S. cerevisiae) homol   Splicing factor, arginine/serine-rich 9   LPS-induced TNF-alpha factor   S755   TGAATACTACT   1,00   0,22   Hs.75867   LPS-induced TNF-alpha factor   S755   TGAAGTCCAGA   1,00   0,22   Hs.75867   LPS-induced TNF-alpha factor   S755   TGAAGTCCAGA   1,00   0,22   Hs.75867   Solute carrier family 20 (phosphate transporter), me   S756   TGTACTTCCTA   1,00   0,22   Hs.75485   Ornithine aminotransferase (gyrate atrophy)   S757   GAGGCAGAAGA   1,00   0,22   Hs.6179   DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k   ESTs, Weakly similar to weak similarity to TPR domai   S759   GAAAGATTGGA   1,00   0,22   Hs.55158   ESTs, Weakly similar to weak similarity to TPR domai   S750   GAAGGCACCAT   1,00   0,22   Hs.279884   DNAJ domain-containing   S761   GAAGGCACCAT   1,00   0,22   Hs.26706   CGI-121 protein   S762   AATGGCACTTA   1,00   0,22   Hs.26706   CGI-121 protein   S763   GATGAGCGGCT   1,00   0,22   Hs.26023   KIAA1041 protein   S765   GCCCAGGGACC   1,00   0,22   Hs.205421   EST   GCCCAGGCACCAT   1	5748	GATGAGTCTCG	10,00			proteasome (prosome, macropain)
5751         TGTTAGCCTGT         1,00         0,22         Hs.92384         vitamin A responsive; cytoskeleton related           5752         TTTTCTGAGTG         1,00         0,22         Hs.82238         POP4 (processing of precursor , S. cerevisiae) homol           5753         ACCAAACTGTG         1,00         0,22         Hs.77608         splicing factor, arginine/serine-rich 9           5754         TGAATACTACT         1,00         0,22         Hs.75867         LPS-induced TNF-alpha factor           5755         TCAAGTCCAGA         1,00         0,22         Hs.75867         solute carrier family 20 (phosphate transporter), me           5756         TGTACTTCCTA         1,00         0,22         Hs.6179         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k           5758         TGCCTATAGTC         1,00         0,22         Hs.55158         ESTs, Weakly similar to weak similarity to TPR domai           5759         GAAAGATTGGA         1,00         0,22         Hs.279884         DNAJ domain-containing           5760         GTCTTCTTAAT         1,00         0,22         Hs.26706         CGI-121 protein           5762         AATGGCACTTA         1,00         0,22         Hs.26023         KIAA1041 protein           5763         GATGACTGCT         1,00         0	5749	ATGGAAAGGAA	2,00	0,44		
related	5750	TGCTGCTTGAA	2,00	0,44	Hs.12152	APMCF1 protein
Cerevisiae   homol	5751	TGTTAGCCTGT	1,00	0,22	Hs.92384	
5754 TGAATACTACT         1,00         0,22 Hs.76507         LPS-induced TNF-alpha factor           5755 TCAAGTCCAGA         1,00         0,22 Hs.75867         solute carrier family 20 (phosphate transporter), me           5756 TGTACTTCCTA         1,00         0,22 Hs.75485         ornithine aminotransferase (gyrate atrophy)           5757 GAGGCAGAAGA         1,00         0,22 Hs.6179         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k           5758 TGCCTATAGTC         1,00         0,22 Hs.55158         ESTs, Weakly similar to weak similarity to TPR domai           5759 GAAAGATTGGA         1,00         0,22 Hs.5076         sorting nexin 12           5760 GTCTTCTTAAT         1,00         0,22 Hs.279884         DNAJ domain-containing           5761 GAAGGCACCAT         1,00         0,22 Hs.26706         CGI-121 protein           5762 AATGGCACTTA         1,00         0,22 Hs.26023         KIAA1041 protein           5763 GATGAGCGGCT         1,00         0,22 Hs.214372         ESTs           5765 GCCCAGGGACC         1,00         0,22 Hs.205421         EST	5752	TTTTCTGAGTG	1,00	0,22	Hs.82238	
5754 TGAATACTACT         1,00         0,22 Hs.76507         LPS-induced TNF-alpha factor           5755 TCAAGTCCAGA         1,00         0,22 Hs.75867         solute carrier family 20 (phosphate transporter), me           5756 TGTACTTCCTA         1,00         0,22 Hs.75485         ornithine aminotransferase (gyrate atrophy)           5757 GAGGCAGAAGA         1,00         0,22 Hs.6179         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k           5758 TGCCTATAGTC         1,00         0,22 Hs.55158         ESTs, Weakly similar to weak similarity to TPR domai           5759 GAAAGATTGGA         1,00         0,22 Hs.5076         sorting nexin 12           5760 GTCTTCTTAAT         1,00         0,22 Hs.279884         DNAJ domain-containing           5761 GAAGGCACCAT         1,00         0,22 Hs.26706         CGI-121 protein           5762 AATGGCACTTA         1,00         0,22 Hs.26023         KIAA1041 protein           5763 GATGAGCGGCT         1,00         0,22 Hs.214372         ESTs           5765 GCCCAGGGACC         1,00         0,22 Hs.205421         EST	5753	ACCAAACTGTG	1,00	0,22	Hs.77608	splicing factor, arginine/serine-rich 9
transporter), me  5756 TGTACTTCCTA  1,00  0,22 Hs.75485  ornithine aminotransferase (gyrate atrophy)  5757 GAGGCAGAAGA  1,00  0,22 Hs.6179  DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k  5758 TGCCTATAGTC  1,00  0,22 Hs.55158  ESTs, Weakly similar to weak similarity to TPR domai  5759 GAAAGATTGGA  1,00  0,22 Hs.5076  sorting nexin 12  5760 GTCTTCTTAAT  1,00  0,22 Hs.279884  DNAJ domain-containing  5761 GAAGGCACCAT  1,00  0,22 Hs.26706  CGI-121 protein  5762 AATGGCACTTA  1,00  0,22 Hs.26023  KIAA1041 protein  5763 GATGAGCGGCT  1,00  0,22 Hs.227637  chromosome 22 open reading frame 2  5764 TGACTGCTGCT  1,00  0,22 Hs.205421  ESTs	5754	TGAATACTACT	1,00	0,22	Hs.76507	
atrophy    5757 GAGGCAGAAGA	5755	TCAAGTCCAGA	1,00	0,22	Hs.75867	
5757         GAGGCAGAAGA         1,00         0,22 Hs.6179         DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 (72k           5758         TGCCTATAGTC         1,00         0,22 Hs.55158         ESTs, Weakly similar to weak similarity to TPR domai           5759         GAAAGATTGGA         1,00         0,22 Hs.5076         sorting nexin 12           5760         GTCTTCTTAAT         1,00         0,22 Hs.279884         DNAJ domain-containing           5761         GAAGGCACCAT         1,00         0,22 Hs.26706         CGI-121 protein           5762         AATGGCACTTA         1,00         0,22 Hs.26023         KIAA1041 protein           5763         GATGAGCGGCT         1,00         0,22 Hs.227637         chromosome 22 open reading frame 2           5764         TGACTGCTGCT         1,00         0,22 Hs.205421         ESTs           5765         GCCCAGGGACC         1,00         0,22 Hs.205421         EST	5756	TGTACTTCCTA	1,00	0,22	Hs.75485	
5758         TGCCTATAGTC         1,00         0,22         Hs.55158         ESTs, Weakly similar to weak similarity to TPR domai           5759         GAAAGATTGGA         1,00         0,22         Hs.5076         sorting nexin 12           5760         GTCTTCTTAAT         1,00         0,22         Hs.279884         DNAJ domain-containing           5761         GAAGGCACCAT         1,00         0,22         Hs.26706         CGI-121 protein           5762         AATGGCACTTA         1,00         0,22         Hs.26023         KIAA1041 protein           5763         GATGAGCGGCT         1,00         0,22         Hs.227637         chromosome 22 open reading frame 2           5764         TGACTGCTGCT         1,00         0,22         Hs.214372         ESTs           5765         GCCCAGGGACC         1,00         0,22         Hs.205421         EST	5757	GAGGCAGAAGA	1,00	0,22	Hs.6179	DEAD/H (Asp-Glu-Ala-Asp/His) box
5760 GTCTTCTTAAT         1,00         0,22 Hs.279884 DNAJ domain-containing           5761 GAAGGCACCAT         1,00         0,22 Hs.26706 CGI-121 protein           5762 AATGGCACTTA         1,00         0,22 Hs.26023 KIAA1041 protein           5763 GATGAGCGGCT         1,00         0,22 Hs.227637 chromosome 22 open reading frame 2           5764 TGACTGCTGCT         1,00         0,22 Hs.214372 ESTs           5765 GCCCAGGGACC         1,00         0,22 Hs.205421 EST	5758	TGCCTATAGTC	1,00	0,22		ESTs, Weakly similar to weak similarity
5760 GTCTTCTTAAT         1,00         0,22 Hs.279884 DNAJ domain-containing           5761 GAAGGCACCAT         1,00         0,22 Hs.26706 CGI-121 protein           5762 AATGGCACTTA         1,00         0,22 Hs.26023 KIAA1041 protein           5763 GATGAGCGGCT         1,00         0,22 Hs.227637 chromosome 22 open reading frame 2           5764 TGACTGCTGCT         1,00         0,22 Hs.214372 ESTs           5765 GCCCAGGGACC         1,00         0,22 Hs.205421 EST	5759	GAAAGATTGGA	1,00	0,22		
5761 GAAGGCACCAT         1,00         0,22 Hs.26706         CGI-121 protein           5762 AATGGCACTTA         1,00         0,22 Hs.26023         KIAA1041 protein           5763 GATGAGCGGCT         1,00         0,22 Hs.227637 chromosome 22 open reading frame 2           5764 TGACTGCTGCT         1,00         0,22 Hs.214372 ESTs           5765 GCCCAGGGACC         1,00         0,22 Hs.205421 EST	5760	GTCTTCTTAAT				
5762 AATGGCACTTA       1,00       0,22 Hs.26023       KIAA1041 protein         5763 GATGAGCGGCT       1,00       0,22 Hs.227637 chromosome 22 open reading frame 2         5764 TGACTGCTGCT       1,00       0,22 Hs.214372 ESTs         5765 GCCCAGGGACC       1,00       0,22 Hs.205421 EST	5761	GAAGGCACCAT				
5763 GATGAGCGGCT         1,00         0,22 Hs.227637 chromosome 22 open reading frame 2           5764 TGACTGCTGCT         1,00         0,22 Hs.214372 ESTs           5765 GCCCAGGGACC         1,00         0,22 Hs.205421 EST	5762	AATGGCACTTA				
5764 TGACTGCTGCT 1,00 0,22 Hs.214372 ESTs 5765 GCCCAGGGACC 1,00 0,22 Hs.205421 EST	5763	GATGAGCGGCT				
5765 GCCCAGGGACC 1,00 0,22 Hs.205421 EST						
	_					
	5766	TAAATGTCCTG	1,00			

<u> </u>					I
5767	GAGAATCTGAT	1,00	0,22	Hs.184352	ESTs, Weakly similar to cDNA EST EMBL:D36107 comes f
5768	TGTTTTGGAAC	1,00	0.22	He 176657	RAN binding protein 3
	TGTAAGTTTTG	1,00			hypothetical protein FLJ20159
	TTGTAGCTCAA	1,00			KIAA0138 gene product
					ESTs
	TGGTCAAGGTG	1,00			
	TATGTGCCACT	1,00	0,22	HS. 1225/9	hypothetical protein FLJ10461
	TATGCCCTATC	1,00			KIAA0210 gene product
	ACTGCTGTCTA	1,00			Homo sapiens HDCMD11P mRNA, partial cds
	AATCTGGTTGC	1,00			heterogeneous nuclear ribonucleoprotein U (scaffold
	AGGAAAGCTGC	54,00			ribosomal protein L36
5777	ACAGTGCTTGC	5,00	0,88	Hs.80350	protein phosphatase 2 (formerly 2A), catalytic subun
5778	GCCCCCCGTG	3,00	0,61	Hs.85573	Homo sapiens mRNA; cDNA
					DKFZp566N034 (from clone DKF
5779	GAGCGGCCTCT	3,00	0,61	Hs.77868	ORF
5780	ACATCGTAGGG	3,00	0,61	Hs.108408	CGI-78 protein
5781	CAAGGGTAAGA	7,00	1,13	Hs.76224	EGF-containing fibulin-like extracellular
			•		matrix pro
5782	TTGTAATCGTG	21,00	2,55	Hs.125078	omithine decarboxylase antizyme 1
5783	GCAAAAAAAA	19,00	2,37		thymosin, beta 10
5784	GGGTGTGTATT	2,00	0,45		angio-associated, migratory cell protein
5785	ATACTGCTGCT	2,00			cullin 2
5786	ACCATAATGTG	2,00		Hs.821	zinc finger protein homologous to Zfp92 in mouse
5787	GACTTGGAGGC	2,00	0,45	Hs.77783	membrane-associated tyrosine- and threonine-specific
5788	GACAGACATCA	2,00	0,45		Homo sapiens mRNA; cDNA
5700	CACCTTCACAC	2.00	0.45	U- 60005	DKFZp434O1230 (from clone DK
	CAGGTTGACAG	2,00			transportin-SR
	GTGCATCCCGA	4,00			casein kinase 2, beta polypeptide
	TOTATACACTT	7,00			H3 histone, family 3A
	TCTATAGAGTT	2,00		Hs.7351	cyclic AMP phosphoprotein, 19 kD
	TTTATTGAAAA	2,00			CD164 antigen, sialomucin
	CGATCAGTTTG	2,00		Hs.34906	
	CTGGGAGGAGG	2,00			transmembrane 4 superfamily member 7
	CTGGTTTCTCC	2,00			p53 inducible protein
	CCTAGGACCTG	2,00			actin related protein 2/3 complex, subunit 4 (20 kD)
5798	AAGGCCACCGG	2,00	0,45	Hs.1384	O-6-methylguanine-DNA methyltransferase
5799	TAAAGTGTCTG	2,00	0,45	Hs.132875	Homo sapiens HSPC309 mRNA, partial cds
1 1					
5800	TTAGGGGGAA	2,00	0,45	Hs.108319	thyroid hormone receptor-associated protein, 150 kDa
	TTTAGGGGGAA TACAATTGTGA	2,00			thyroid hormone receptor-associated protein, 150 kDa hypothetical protein DKFZp586H0623

E000	CCTCCCCTCCC	4.00	0.77	Ha 02070	ECTs Wookly similar to missatubula
	CGTGGGGTGGC	4,00		Hs.92679	ESTs, Weakly similar to microtubule-based motor [H.s
	AGCTTGGACAG	1,00		Hs.92700	DKFZP564O243 protein
5805	TATGTTCAGAA	1,00	0,23	Hs.6241	phosphoinositide-3-kinase, regulatory subunit, polyp
5806	CTACAATAAAT	1,00	0,23	Hs.57100	ESTs
5807	TATCCTGGTAA	1,00	0,23	Hs.284291	sorting nexin 6
5808	CAAATAAACTT	1,00			ESTs
5809	CTGCCCTGGGA	1,00			ribosomal protein S3
5810	CCAGAAAGAAT	1,00			tissue inhibitor of metalloproteinase 3 (Sorsby fund
	ATCTCTTTCCC	1,00			ESTs, Highly similar to serine protease [H.sapiens]
	TCTGTTATTGG	1,00			hypothetical protein FLJ10024
	GAGCACATCAG	1,00	0,23	Hs.173497	Sec23 (S. cerevisiae) homolog B
	CCATCCAGTGA	1,00	0,23	Hs.155140	casein kinase 2, alpha 1 polypeptide
	GCCGGCCCGGC	1,00	0,23	Hs.133230	ribosomal protein S15
5816	CAAACCATCCG	1,00	0,23	Hs.125170	ESTs
5817	TTCTCTACACA	5,00	0,91		transforming growth factor beta- stimulated protein T
5818	GAGGGTGGCGC	3,00	0,63	Hs.81687	non-metastatic cells 3, protein expressed in
5819	TTGTCCAGAGG	3,00	0,63	Hs.14839	polymerase (RNA) II (DNA directed) polypeptide G
5820	CTTGTAATCCT	7,00	1,16	Hs.265827	interferon, alpha-inducible protein (clone IFI-6-16)
5821	GGCTGATTTTT	2,00	0,46	Hs.81361	heterogeneous nuclear ribonucleoprotein A/B
5822	TGCTCTGTGTA	2,00	0,46	Hs.70186	suppressor of Ty (S.cerevisiae) 5 homolog
5823	TATGTAATATG	2,00	0,46	Hs.41569	phosphatidic acid phosphatase type 2A
5824	CCACAACCTGG	2,00	0,46	Hs.101742	Human DNA sequence from clone 321D2 on chromosome 16
5825	GAAAAATGGTT	49,00	5,38	Hs.283106	hemoglobin, gamma A
5826	ACGGAACAATA	8,00	1,29	Hs.8272	prostaglandin D2 synthase (21kD, brain)
5827	TAAAAGACAAA	3,00	0,63	Hs.77196	spectrin, alpha, non-erythrocytic 1 (alpha-fodrin)
5828	CTCAACATCTC	44,00	4,98		ribosomal protein, large, P0
5829	CAATCACAAAA	2,00			calpain 2, (m/ll) large subunit
5830	AATGCAAGATA	2,00			transcription elongation factor B (SIII), polypeptid
5831	CCCCTCCCCAG	2,00	0,46	Hs.150540	Homo sapiens chromosome 22q13 BAC clone CIT987SK-384
5832	AGCTCTCCCTG	48,00	5,42	Hs.82202	ribosomal protein L17
5833	GATGTCTCTAG	3,00		Hs.74368	transmembrane protein (63kD),
					endoplasmic reticulum/
5834	AATAAAGTTGT	4,00	0,79	Hs.117176	poly(A)-binding protein, nuclear 1
	TGGCCAAAAAA	1,00			ESTs
5836	AATAGCTGATT	1,00	0,24	Hs.90011	adenylosuccinate synthase

5838 GAGGAAGGCTC         1,00         0,24 Hs.78771         phosphoglycerate kinase 1           5839 CTATTAGGGA         1,00         0,24 Hs.7736         hypothetical protein           5840 CTAGCAGAGCA         1,00         0,24 Hs.4747         dyskeratosis congenita 1, dyskerin           5841 GCCTGTGGGTT         1,00         0,24 Hs.43549         uncharacterized hematopoietic stem/progenitor cells           5842 TTTTGGATGTA         1,00         0,24 Hs.182874         Homo sapiens ubiquitin-conjugatin enzyme variant Ku           5843 ACCGGGGTGAA         1,00         0,24 Hs.182874         guanine nucleotide binding protein protein) alpha           5845 AAGACAGAGCA         1,00         0,24 Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5846 GGGCAGAATAT         1,00         0,24 Hs.168670 peroxisomal farnesylated protein protein [H.sapi           5848 GACAAGCCGG         5,00         0,93 Hs.155433 ATP synthase, H+ transporting, mitochondrial F1 comp           5849 AGACAAGCTGG         4,00         0,80 Hs.168975 splicing factor, arginine/serine-rich serine-rich serine-	
5840 CTAGCAGAGCA         1,00         0,24 Hs.4747         dyskeratosis congenita 1, dyskerin           5841 GCCTGTGGGTT         1,00         0,24 Hs.43549         uncharacterized hematopoietic stem/progenitor cells           5842 TTTTGGATGTA         1,00         0,24 Hs.286047         Homo sapiens ubiquitin-conjugatin enzyme variant Ku           5843 ACCGGGGTGAA         1,00         0,24 Hs.182874         guanine nucleotide binding protein protein jalpha           5845 AAGACAGAGCA         1,00         0,24 Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5846 GGCAGAATAT         1,00         0,24 Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5847 TGATCGCGGCT         1,00         0,24 Hs.135906         ESTs           5848 GACAAGCCAGG         5,00         0,93 Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5849 AGACAAGCTGG         4,00         0,80 Hs.166975         splicing factor, arginine/serine-rich           5850 GCTGGTGCCTG         4,00         0,80 Hs.125359         Thy-1 cell surface antigen           5851 GCTAAAAAAAA         3,00         0,64 Hs.256697         histline triad nucleotide-binding protein 2 (49kD           5853 CTGGGTCTCCA         4,00         0,80 Hs.180842 ribosomal protein L13           5855 GAAAAAATGTT         2,00         0,47 Hs.194329	
5841         GCCTGTGGGTT         1,00         0,24         Hs.43549         uncharacterized hematopoietic stern/progenitor cells           5842         TTTTGGATGTA         1,00         0,24         Hs.286047         Homo sapiens ubiquitin-conjugatin enzyme variant Ku           5843         ACCGGGGTGAA         1,00         0,24         Hs.182874         guanine nucleotide binding protein protein alpha           5844         GCCCCGGAGCC         1,00         0,24         Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5845         AAGACAGAGCA         1,00         0,24         Hs.168670 peroxisomal farnesylated protein           5846         GGGCAGAATAT         1,00         0,24         Hs.135906         ESTs, Weakly similar to hypothetic protein [H.sapi           5847         TGATCGCGGCT         1,00         0,24         Hs.135906         ESTs           5849         AGACAAGCCTGG         4,00         0,80         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5850         GCTGGTGCCTG         4,00         0,80         Hs.125359         Thy-1 cell surface antigen           5851         GCTGGTGCTG         4,00         0,80         Hs.173611         NADH dehydrogenase (ubiquinone protein 2 (49kD           5852         TACTAAAAAAA	
Stem/progenitor cells	
5842         TTTTGGATGTA         1,00         0,24         Hs.286047         Homo sapiens ubiquitin-conjugatinenzyme variant Ku           5843         ACCGGGGTGAA         1,00         0,24         Hs.182874         guanine nucleotide binding protein protein alpha           5844         GCCCCGGAGCC         1,00         0,24         Hs.17528         amyloid beta (A4) precursor protein binding, family           5845         AAGACAGAGCA         1,00         0,24         Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5846         GGGCAGAATAT         1,00         0,24         Hs.168670         peroxisomal farnesylated protein           5847         TGATCGCGGCT         1,00         0,24         Hs.15906         ESTs           5849         GACAATGCCAG         5,00         0,93         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp mitochondrial F1 comp mitochondrial F1 comp splicing factor, arginine/serine-rich           5850         GCTGGTGCCTG         4,00         0,80         Hs.125359         Thy-1 cell surface antigen           5851         GCTAAAAAAAA         3,00         0,64         Hs.173611         NADH dehydrogenase (ubiquinone protein) associated           5853         CTGGGTCTCCA         4,00         0,80         Hs.180842         ribosomal protein L13	
enzyme variant Ku	
5843         ACCGGGGTGAA         1,00         0,24         Hs.182874         guanine nucleotide binding protein protein) alpha           5844         GCCCCGGAGCC         1,00         0,24         Hs.17528         amyloid beta (A4) precursor protein binding, family binding, family           5845         AAGACAGAGCA         1,00         0,24         Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5846         GGGCAGAATAT         1,00         0,24         Hs.155906         ESTs           5847         TGATCGCGGCT         1,00         0,24         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5848         GACAAGCTGG         4,00         0,80         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5849         AGACAAGCTGG         4,00         0,80         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5850         GCTGGTGCTG         4,00         0,80         Hs.125359         Thy-1 cell surface antigen           5851         GCTAAAAAAAA         3,00         0,64         Hs.173611         NADH dehydrogenase (ubiquinone protein 2 (49kD           5853         CTGGGTCTCCA         4,00         0,80         Hs.180842         ribosomal protein L13           5854	3
5844         GCCCCGGAGCC         1,00         0,24         Hs.17528         amyloid beta (A4) precursor protein binding, family           5845         AAGACAGAGCA         1,00         0,24         Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi           5846         GGGCAGAATAT         1,00         0,24         Hs.168670         peroxisomal farnesylated protein           5847         TGATCGCGGCT         1,00         0,24         Hs.135906         ESTs           5848         GACAATGCCAG         5,00         0,93         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5849         AGACAAGCTGG         4,00         0,80         Hs.166975         splicing factor, arginine/serine-rich           5850         GCTGGTGCCTG         4,00         0,80         Hs.125359         Thy-1 cell surface antigen           5851         GCTAAAAAAAA         3,00         0,64         Hs.256697         histidine triad nucleotide-binding protein           5852         TACTAAAAAAAA         3,00         0,64         Hs.173611         NADH dehydrogenase (ubiquinone protein 2 (49kD           5853         CTGGGTCTCCA         4,00         0,80         Hs.180842         ribosomal protein L13           5854         GTATTTAACAT         2,00 <td< td=""><td>(G</td></td<>	(G
5845         AAGACAGAGCA         1,00         0,24         Hs.169854         ESTs, Weakly similar to hypothetic protein [H.sapi]           5846         GGGCAGAATAT         1,00         0,24         Hs.168670         peroxisomal farnesylated protein           5847         TGATCGCGGCT         1,00         0,24         Hs.135906         ESTs           5848         GACAATGCCAG         5,00         0,93         Hs.155433         ATP synthase, H+ transporting, mitochondrial F1 comp           5849         AGACAAGCTGG         4,00         0,80         Hs.166975         splicing factor, arginine/serine-rich splicing factor, arginin	}-
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5847 TGATCGCGGCT         1,00         0,24 Hs.135906 ESTs           5848 GACAATGCCAG         5,00         0,93 Hs.155433 ATP synthase, H+ transporting, mitochondrial F1 comp           5849 AGACAAGCTGG         4,00         0,80 Hs.166975 splicing factor, arginine/serine-rich           5850 GCTGGTCCTG         4,00         0,80 Hs.125359 Thy-1 cell surface antigen           5851 GCTAAAAAAAA         3,00         0,64 Hs.256697 histidine triad nucleotide-binding protein L13           5852 TACTAAAAAAA         3,00         0,64 Hs.173611 NADH dehydrogenase (ubiquinone protein 2 (49kD)           5853 CTGGGTCTCCA         4,00         0,80 Hs.180842 ribosomal protein L13           5854 GTATTTAACAT         2,00         0,47 Hs.194329 ESTs, Weakly similar to AF125535 pp21 homolog [H.s           5855 GAAAAAATGTT         2,00         0,47 Hs.189583 putative protein similar to nessy (Drosophila)           5857 TCAAGAAATTA         2,00         0,47 Hs.152978 proteasome (prosome, macropain) activator subunit 3           5858 GAATCCGATTT         2,00         0,47 Hs.151536 RAB13, member RAS oncogene factivator subunit 3           5859 GCGCTTCTAG         3,00         0,65 Hs.75813 polycystic kidney disease 1 (autost dominant)           5860 AGCGTGGATG         2,00         0,47 Hs.116449 ESTs, Weakly similar to AF123303 calcium-binding t           5861 GAGGACAAAATA         2,00         0,47 Hs.116449 ESTs	
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5854 GTATTTAACAT  2,00  0,47 Hs.9006  VAMP (vesicle-associated membra protein)-associate  5855 GAAAAAATGTT  2,00  0,47 Hs.194329 ESTs, Weakly similar to AF125535 pp21 homolog [H.s  5856 GGGGGGTGGAG  2,00  0,47 Hs.189583 putative protein similar to nessy (Drosophila)  5857 TCAAGAAATTA  2,00  0,47 Hs.152978 proteasome (prosome, macropain) activator subunit 3  5858 GAATCCGATTT  2,00  0,47 Hs.151536 RAB13, member RAS oncogene fa  5859 GCCGCTTCTAG  3,00  0,65 Hs.75813 polycystic kidney disease 1 (autosi dominant)  5860 AGCGTGTGATG  2,00  0,47 Hs.74649 cytochrome c oxidase subunit VIc  5861 GAGGGACTTGG  2,00  0,47 Hs.32246 ESTs, Weakly similar to AF123303 calcium-binding t  5862 CAAGCAAAATA  2,00  0,47 Hs.116449 ESTs  5863 AGGAGCAAAGG  5,00  0,96 Hs.76289 biliverdin reductase B (flavin reductase B)  5864 GCCCTGACCTT  1,00  0,24 Hs.97871 ESTs, Moderately similar to hypother	
5855 GAAAAAATGTT  2,00  0,47 Hs.194329 ESTs, Weakly similar to AF125535 pp21 homolog [H.s  5856 GGGGGGTGGAG  2,00  0,47 Hs.189583 putative protein similar to nessy (Drosophila)  5857 TCAAGAAATTA  2,00  0,47 Hs.152978 proteasome (prosome, macropain) activator subunit 3  5858 GAATCCGATTT  2,00  0,47 Hs.151536 RAB13, member RAS oncogene fa polycystic kidney disease 1 (autose dominant)  5860 AGCGTGTGATG  2,00  0,47 Hs.74649 cytochrome c oxidase subunit Vic ESTs, Weakly similar to AF123303 calcium-binding t  5862 CAAGCAAAATA  2,00  0,47 Hs.116449 ESTs  5863 AGGAGCAAAGG  5,00  0,96 Hs.76289 biliverdin reductase B (flavin reductase B)  (NADPH))  5864 GCCCTGACCTT  1,00  0,24 Hs.97871 ESTs, Moderately similar to hypothesis and the polycystic kidney disease 1 (autose dominant)  5865 CCCTGACCTT  1,00  0,47 Hs.116449 ESTs  5866 GCCCTGACCTT  1,00  0,24 Hs.97871 ESTs, Moderately similar to hypothesis and the polycystic kidney disease 2 (autose dominant)  5867 CAAGCAAAATA  2,00  0,47 Hs.116449 ESTs  1,00  1,	ne
5856GGGGGTGGAG2,000,47Hs.189583putative protein similar to nessy (Drosophila)5857TCAAGAAATTA2,000,47Hs.152978proteasome (prosome, macropain) activator subunit 35858GAATCCGATTT2,000,47Hs.151536RAB13, member RAS oncogene fa5859GCCGCTTCTAG3,000,65Hs.75813polycystic kidney disease 1 (autosidominant)5860AGCGTGTGATG2,000,47Hs.74649cytochrome c oxidase subunit VIc5861GAGGGACTTGG2,000,47Hs.32246ESTs, Weakly similar to AF123303 calcium-binding t5862CAAGCAAAATA2,000,47Hs.116449ESTs5863AGGAGCAAAGG5,000,96Hs.76289biliverdin reductase B (flavin reductase B)5864GCCCTGACCTT1,000,24Hs.97871ESTs, Moderately similar to hypothesical contents of the contents of	_1
5857 TCAAGAAATTA  2,00  0,47 Hs.152978 proteasome (prosome, macropain) activator subunit 3  5858 GAATCCGATTT  2,00  0,47 Hs.151536 RAB13, member RAS oncogene fa polycystic kidney disease 1 (autosidominant)  5860 AGCGTGTGATG  2,00  0,47 Hs.74649 cytochrome c oxidase subunit VIc  5861 GAGGGACTTGG  2,00  0,47 Hs.32246 ESTs, Weakly similar to AF123303 calcium-binding t  5862 CAAGCAAAATA  2,00  0,47 Hs.116449 ESTs  5863 AGGAGCAAAGG  5,00  0,96 Hs.76289 biliverdin reductase B (flavin reductase B) (NADPH))  5864 GCCCTGACCTT  1,00  0,24 Hs.97871 ESTs, Moderately similar to hypothesis and the company of the	
5858 GAATCCGATTT2,000,47 Hs.151536 RAB13, member RAS oncogene fa5859 GCCGCTTCTAG3,000,65 Hs.75813 polycystic kidney disease 1 (autosidominant)5860 AGCGTGTGATG2,000,47 Hs.74649 cytochrome c oxidase subunit VIc5861 GAGGGACTTGG2,000,47 Hs.32246 ESTs, Weakly similar to AF123303 calcium-binding t5862 CAAGCAAAATA2,000,47 Hs.116449 ESTs5863 AGGAGCAAAGG5,000,96 Hs.76289 biliverdin reductase B (flavin reductase B)5864 GCCCTGACCTT1,000,24 Hs.97871 ESTs, Moderately similar to hypothesis	
5859GCCGCTTCTAG3,000,65Hs.75813polycystic kidney disease 1 (autosidominant)5860AGCGTGTGATG2,000,47Hs.74649cytochrome c oxidase subunit VIc5861GAGGGACTTGG2,000,47Hs.32246ESTs, Weakly similar to AF123303 calcium-binding t5862CAAGCAAAATA2,000,47Hs.116449ESTs5863AGGAGCAAAGG5,000,96Hs.76289biliverdin reductase B (flavin reductase B)5864GCCCTGACCTT1,000,24Hs.97871ESTs, Moderately similar to hypothesis	mily
5860 AGCGTGTGATG2,000,47 Hs.74649cytochrome c oxidase subunit VIc5861 GAGGACTTGG2,000,47 Hs.32246ESTs, Weakly similar to AF123303 calcium-binding t5862 CAAGCAAAATA2,000,47 Hs.116449 ESTs5863 AGGAGCAAAGG5,000,96 Hs.76289biliverdin reductase B (flavin reduc (NADPH))5864 GCCCTGACCTT1,000,24 Hs.97871ESTs, Moderately similar to hypoth	
5861 GAGGGACTTGG   2,00   0,47 Hs.32246   ESTs, Weakly similar to AF123303 calcium-binding t   5862 CAAGCAAAATA   2,00   0,47 Hs.116449 ESTs   5863 AGGAGCAAAGG   5,00   0,96 Hs.76289   biliverdin reductase B (flavin reductable (NADPH))   5864 GCCCTGACCTT   1,00   0,24 Hs.97871   ESTs, Moderately similar to hypothesis   1,00   1	
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5863 AGGAGCAAAGG 5,00 0,96 Hs.76289 biliverdin reductase B (flavin	
5864 GCCCTGACCTT 1,00 0,24 Hs.97871 ESTs, Moderately similar to hypoth	ase
	etical
5865 AGTAAAACTTC 1,00 0,24 Hs.9552 binder of Arl Two	
5866 TCTAAAGAATG 1,00 0,24 Hs.92918 hypothetical protein	
5867 TATCTTCTAAG 1,00 0,24 Hs.82314 hypoxanthine phosphoribosyltrans 1 (Lesch-Nyha	erase
5868 GTTCTATTTAA 1,00 0,24 Hs.8033 hypothetical protein FLJ11164	

5060	CCACCCACTCT	1.00	0.24	Ha 74040	lauta abrama a avida a a cubumit \//la
	CCAGGCACTGT	1,00		Hs.74649	cytochrome c oxidase subunit VIc
	GCTGCCTGCCT	1,00		Hs.71962	ESTs
	CTAAAACTGGT	1,00		Hs.69993	ESTs Control of the c
	AACCTGGCCTC	1,00		Hs.6780	protein tyrosine kinase 9-like (A6-related protein)
5873	TACACCAAGAA	1,00	0,24	Hs.6449	hypothetical protein FLJ20542
<u></u>	GAGCTGCAGGG	. 1,00		Hs.6294	hypothetical protein DKFZp434L1435 similar to valyl
	TGTATATGGTG	1,00	0,24	Hs.279849	KIAA0438 gene product
5876	CTTGAGTCACA	1,00	0,24	Hs.261023	Human DNA sequence from clone RP3-500L14 on chromoso
5877	CTTTGTTTTGC	1,00	0,24	Hs.2271	endothelin 1
5878	GATTTAAAAAA	1,00	0,24	Hs.187991	DKFZP564A122 protein
5879	TGATGATGTTG	1,00			F-box only protein 9
5880	CTGGCTCCATC	1,00		Hs.10290	U5 snRNP-specific 40 kDa protein (hPrp8-binding)
5881	CTCACACATTA	5,00		Hs.150101	lysosomal-associated membrane protein
5882	GGGCCAGGGGA	5,00	0,97	Hs.111099	ESTs, Weakly similar to alpha-1 type I collagen [H.s
5883	GGCCCGGCTTT	2,00	0,48	Hs.283639	ESTs
5884	TGAGGCAGGGA	2,00	0,48	Hs.154546	syntaxin 5A
5885	GGCAAGAAGAA	27,00			ribosomal protein L27
5886	CATCTAAACTG	7,00			Williams-Beuren syndrome chromosome region 1
5887	TGTTCATCATC	6,00	1,12	Hs.65450	
5888	CAGCAGAAGCA	14,00			small EDRK-rich factor 2
5889	TTGAATTCCCC	6,00			sema domain, immunoglobulin domain (Ig), short basic
5890	ACAGTCTTGCC	2,00	0,48	Hs.77665	KIAA0102 gene product
5891	AGCACTTTTGC	2,00	0,48	Hs.103419	fasciculation and elongation protein zeta 2 (zygin I
5892	GAGCAGCGCCC	7,00	1,27	Hs.112408	S100 calcium-binding protein A7 (psoriasin 1)
	AGTTTGGGCTG	1,00		Hs.9911	ESTs
	GTGTTGAGAGA	1,00			methionine adenosyltransferase II, alpha
5895	CCCCTGCTGCT	1,00	0,25	Hs.6272	Homo sapiens mRNA; cDNA DKFZp761P0114 (from clone DK
5896	GCCTGTTTGTA	1,00	0,25	Hs.5613	Homo sapiens mRNA; cDNA DKFZp564E2222 (from clone DK
5897	TACGCTTGGTC	1,00	0.25	Hs.284271	cytochrome b5 reductase 1 (B5R.1)
	TGGGGATTACC	1,00			Homo sapiens unknown mRNA
	TGGTTTATTAA	1,00		Hs.23023	
	TCCAGGGCTCT	1,00			MYB binding protein (P160) 1a
	CCAGTAGAAGT	1,00			HSPC039 protein
	TGGTTTTGTAT	1,00			KIAA0175 gene product
	TAAATTITAAA	1,00			cofilin 1 (non-muscle)
	GTGTGGGAGAT	1,00		Hs.16331	
	GAAGTGCCCAT	1,00			KIAA1237 protein
			<u> </u>	. 10. 10 70 1	i watter brotoni

5006	CTAGCCTCACG	48,00	6.03	Hs.14376	actin, gamma 1
	TGCCTGCTCCT	3,00		Hs.93678	ESTs
	GCCGCCTGCCT	2,00		Hs.850	
13900	GCCGCCTGCCT	2,00	0,49	ns.650	IMP (inosine monophosphate) dehydrogenase 1
Sana	AGCCGGGCTTT	2,00	0.40	Hs.57079	ESTs
	GAGGGGATGTA	2,00			RAB31, member RAS oncogene family
	ACTITGAATGA	2,00		Hs.19077	
	CTGCCCGCCTG			Hs.15961	CGI-113 protein
	CTGTTGGTGAT	2,00			dynactin 3 (p22)
_	TGTGCTCAGGG	38,00		Hs.3463	ribosomal protein S23
3914	I G I G C I C A G G G	1,00	0,25	Hs.92261	pyruvate dehydrogenase kinase,
5015	TTCTTTGGGAA	1,00	0.25	Hs.9095	isoenzyme 2 Homo sapiens mRNA; cDNA
3313	1101110000	1,00	0,23	ns.9095	DKFZp434N1710 (from clone DK
5916	GGCATCAGGGG	1,00	0.25	Hs.8361	hypothetical protein, estradiol-induced
	TCTGCAAGAAG	1,00		Hs.81281	hypothetical protein
	CGTGGCCACGG	1,00		Hs.77221	choline kinase
	GTGTGCTTAGA	1,00			
	GTGGGGAGGAC	1,00		Hs.7165	hypothetical protein FLJ20038
	TCTGTCAATCT			Hs.65234	zinc finger protein 259
	AATAAACGTGT	1,00		Hs.57304	hypothetical protein FLJ20596
	AGCGCCTTCCT	1,00			Ras-related GTP-binding protein
	CTCAGCCTGAA	1,00		Hs.49005 Hs.3496	hypothetical protein
	GCAGTGCCACT	1,00			ESTs
	CTAGAAGTACA	1,00			ESTs
		1,00		Hs.32500	acetyl-Coenzyme A acyltransferase 2 (mitochondrial 3
	ATTITGTCGTG	1,00			hypothetical protein FLJ11015
	GATTTCAGCTC	1,00	0,25	Hs.280978	hypothetical protein FLJ10496
	TCACAGCAAGG	1,00	0,25	Hs.279944	replication initiation region protein (60kD)
· ·	GCTGGGCGGCT	1,00	0,25	Hs.272788	Homo sapiens cDNA FLJ20205 fis, clone COLF1506
5931	AGGATATCCAG	1,00	0,25		pleiotropic regulator 1 (PRL1, Arabidopsis homolog)
5932	TATTCACTAAA	1,00	0.25		trinucleotide repeat containing 3
5933	GGCCTCCAGCC	1,00	0.25	Hs.184669	zinc finger protein 144 (Mel-18)
5934	ATTTGTATCTA	1,00	0.25	Hs.181363	ESTs
5935	AGTAAGCTCCC	1,00			ALEX3 protein
5936	CCCTCGCATTG	1,00			Human DNA sequence from clone
			, , ,		321D2 on chromosome 16
5937	GAGAGAAGAGT	1,00	0,25		UDP-Gal:betaGlcNAc beta 1,4-
			<i>'</i>		galactosyltransferase,
5938	GATTACTTGCC	1,00	0,25		peroxisomal biogenesis factor 16
5939	TTCTTATTTTA	3,00			splicing factor 3b, subunit 2, 145kD
5940	GGGTCAAAAGG	8,00			H3 histone, family 3A
	GAGTTGGCAGT	4,00			heme-regulated initiation factor 2-alpha
}		•	. 1		kinase
5942	GCCTTTCCCTC	4,00	0,86		ubiquitin specific protease 22
	GCATATGAGCA	2,00		Hs.7149	ESTs, Weakly similar to hypothetical
					protein [H.sapi
5944	AACAGAATATG	2,00	0,50	Hs.69469	dendritic cell protein

EQ4E	TGTGCTAAATG	27.00	2 02	U- 250005	with a general protoin 1.24
		27,00			ribosomal protein L34
	GAAATGTAAGA	9,00			poly(rC)-binding protein 2
	TGAAGAGAATT	3,00			
5948	CTGCTTAAGGG	2,00	0,50	Hs.76662	ESTs, Weakly similar to KIAA0946
					protein [H.sapiens]
	GCTCAGGTCTG	2,00			endothelin converting enzyme 1
	TTCTTGCTTAA	2,00			ubiquitin-conjugating enzyme E2L 6
5951	TTTCTTAAAGG	4,00	0,88	Hs.197114	RNA binding protein; AT-rich element binding factor
5952	TAGAATGCAAA	1,00	0.26	Hs.7946	KIAA1288 protein
	CTTCCCACTCT	1,00		Hs.6845	ESTs, Weakly similar to JC4903 UDP-
		.,00			galactose transpo
5954	AAATCCTAGAA	1,00	0,26	Hs.6592	ESTs, Weakly similar to salivary prolinerich protei
	ACAGCTAATTA	1,00			eukaryotic translation initiation factor 1A
5956	TTTCATCTGTA	1,00	0,26	Hs.285013	putative human HLA class II associated
E057	CCTCTTCACAC	4.00	0.00	11- 004400	protein I
	CGTGTTGAGAG	1,00			PRO2047 protein
5958	TCTACTGTTAG	1,00	0,26		Human DNA sequence from clone RP3-460J8 on chromosom
5959	TAAATAAAGCA	1,00	0,26	Hs.214783	ESTs
5960	AACTGGGTCTG	1,00	0,26	Hs.182215	ADP-ribosylation factor-like 3
	AATACATCAAA	1,00		Hs.178761	26S proteasome-associated pad1 homolog
5962	CCTGAGTTGAT	1,00	0.26		clone FLB4739
	AAATCTCTGGT	1,00			KIAA1415 protein
	TGCTACGATCT	1,00		Hs.103135	
-	GCGACAGCTCC	9,00			ribosomal protein L24
$\overline{}$	ATACAATAAAA	3,00			nuclear transport factor 2 (placental
	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	3,00	0,7 1	1 13. 13 17 37	protein 15)
5967	GAGGCCGACCC	2,00	0,51	Hs.74631	basigin
5968	GCAAGACCCCA	2,00			KIAA0562 gene product
	GGCCGCTGCTC	2,00	0,51	Hs.151531	protein phosphatase 3 (formerly 2B), catalytic subun
	GGTGAGACACT	12,00	2,05	Hs.164280	solute carrier family 25 (mitochondrial carrier; ade
5971	TCTGTTTATCA	7,00	1,36	Hs.180394	signal recognition particle 14kD (homologous Alu RNA
5972	TGCTGGGTGGG	4,00	0,89	Hs.198273	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 8
5973	TCACCTGTAGG	2,00	0.51	Hs.78921	A kinase (PRKA) anchor protein 1
	CTTGACACACA	2,00			eukaryotic translation initiation factor 5
	TGAAGGAGCCG	12,00			ATP synthase, H+ transporting,
		,	,		mitochondrial F0 comp
5976	CTTTGATGTTC	3,00	0,72	Hs.85100	WD repeat domain 1
	TCGTCTTTATC	13,00			ribosomal protein S7
	ATGGCAAGGGA	7,00			DR1-associated protein 1 (negative cofactor 2 alpha)
5979	AGTAAAAAAA	1,00	0.26	Hs.9552	binder of Arl Two
3919	V01VVVVVVV	1,00	0,20	115.5002	Dilitidat Of WILLIMO

5980	CAGTGGAATGC	1,00		Hs.811	ubiquitin-conjugating enzyme E2B (RAD6 homolog)
5981	GTGTCTTGTAG	1,00	0,26	Hs.6166	hypothetical protein FLJ11196
5982	CAAAAGGCTCT	1,00	0,26	Hs.5392	potassium channel modulatory factor
5983	TATAGTGGCTA	1,00	0,26	Hs.5326	Homo sapiens SSX4 protein gene, partial cds; OATL1 p
5984	GTTCTCTTTGT	1,00	0,26	Hs.285458	Homo sapiens mRNA; cDNA DKFZp434G012 (from clone DKF
5985	CACAAGCTTCA	1,00	0,26	Hs.284176	hypothetical protein PRO2221
5986	ACGTGAGTGCT	1,00			CGI-105 protein
5987	TCTTCTGCTTT	1,00	0,26		hepatocellular carcinoma-associated antigen 59
5988	CCTTCCCTGAG	1,00	0,26	Hs.26584	diaphanous (Drosophila, homolog) 1
5989	CCTGGAGGGGC	1,00		Hs.265327	Human DNA sequence from clone RP4-756G23 on chromoso
5990	TGGATGCTGTT	1,00	0.26		hypothetical protein FLJ10604
	CAGCCCTCCCG	1,00		Hs.250857	calcium/calmodulin-dependent protein kinase (CaM kin
5992	TATGGTACCAA	1,00	0.26	Hs.242710	
$\overline{}$	AGAGACTCTTG	1,00			DKFZP547E1010 protein
5994	CAGCCATAAAA	1,00			hypothetical protein FLJ20724
5995	AAGCTGTATAA	1,00			hexabrachion (tenascin C, cytotactin)
	TGGCCTAATAT	1,00		Hs.1501	syndecan 2 (heparan sulfate proteoglycan 1, cell sur
5997	TTGCTGTAGAC	1,00	0.26		p53 regulated PA26 nuclear protein
	TTACTGCCTAG	1,00			ESTs
	TAGCTGCCTTT	1,00			CGI-43 protein
	CTGGGTTGTGA	1,00			KIAA1249 protein
6001	TGGTTTGCGTA	3,00	0,73	Hs.6459	ESTs, Moderately similar to unnamed protein product
	CTCATCAGCTT	5,00	1,08	Hs.104125	adenylyl cyclase-associated protein
	CACCCCCAGGC	2,00	0,52	Hs.7301	G protein pathway suppressor 2
	CAAAAAAAAA	14,00	2,41	Hs.112272	histone deacetylase 8
	CCGCCGAAGTT	5,00	1,08	Hs.182979	ribosomal protein L12
	TGTACCTGTAA	33,00			tubulin, alpha, ubiquitous
	AAAAAAAAAAG	3,00			ribosomal protein L13
6008	TGGATCAACCA	4,00	0,92	Hs.286030	caveolin 1 caveolae protein, 22kD
	GAAGTGTGTCC	2,00			CGI-45 protein
	CCCAATTTTCA	2,00			polymerase (RNA) II (DNA directed) polypeptide F
6011	CAGATTTGCAT	2,00	0,53	Hs.155986	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 24
6012	GGTCCCCTACC	2,00	0,53	Hs.151761	KIAA0100 gene product
	AATCCTGTGGA	39,00			ribosomal protein L8
	AGGGGATTCCC	3,00			Arginine-rich protein
	CTTTCTTTGAG	5,00		Hs.4909	dickkopf (Xenopus laevis) homolog 3
	AGAGACAAGTC	2,00		Hs.98614	ribosome binding protein 1 (dog 180kD homolog)
6017	TGTTTTCGCCT	2,00	0,53	Hs.83916	NADH dehydrogenase (ubiquinone) 1

	Γ			Γ	alpha subcomplex,
6018	GGCCCACACCC	2,00	0.53	Hs.77978	Homo sapiens HRIHFB2007 mRNA,
					partial cds
6019	GTGAAACATTA	1,00	0,27	Hs.30352	ribosomal protein S6 kinase, 52kD, polypeptide 1
6020	GATTTGAAATG	1,00	0,27	Hs.273307	signal recognition particle 68kD
	CACAGTCAAAA	1,00			calcium channel, voltage-dependent, beta 3 subunit
6022	TAACCCAACAA	1,00		Hs.1869	phosphoglucomutase 1
6023	TGGGAAAACTC	1,00	.0,27	Hs.153028	cytochrome b-561
6024	AAATCGATGAG	1,00	0,27	Hs.13580	ESTs, Weakly similar to AF180920_1 cyclin L ania-6a
6025	GGAAGATGAAC	1,00	0,27	Hs.12255	hypothetical protein FLJ10656
6026	AGGTTTTGCCT	1,00	0,27	Hs.102469	putative nuclear protein
6027	GCCCTCGGCCG	4,00	0,93	Hs.17883	protein phosphatase 1G (formerly 2C), magnesium-depe
6028	CGGAGACCCTA	3,00	0,74	Hs.74497	nuclease sensitive element binding protein 1
6029	AATGAAAAAA	3,00		Hs.11393	RAD51 (S. cerevisiae) homolog C
6030	CTGCTATACGA	17,00	2,94	Hs.180946	ribosomal protein L5
6031	CCACTCCTCAA	7,00	1,45	Hs.82890	defender against cell death 1
6032	CCCAACCCCTG	3,00	0,75	Hs.169488	dentatorubral-pallidoluysian atrophy (atrophin-1)
6033	CCTGAGCCCGG	5,00	1,12	Hs.68571	VPS28 protein
6034	GCGCTGGAGTG	7,00			ESTs, Weakly similar to B0495.6 [C.elegans]
6035	TGGCAGCTTTT	3,00	0,76	Hs.6153	CGI-48 protein
6036	CAGCTTCACCT	2,00	0,55	Hs.179606	nuclear RNA helicase, DECD variant of DEAD box famil
6037	AGTGGCTGTGT	1,00	0,28	Hs.5486	clone FLB5214
6038	ACTCCTTCCTC	1,00			S100 calcium-binding protein A11 (calgizzarin)
6039	GCAAATCTGAA	1,00	0,28	Hs.242894	ADP-ribosylation factor-like 1
6040	TAAAGGTTTTT	1,00	0,28	Hs.241520	transcriptional coactivator
6041	CCTTTTGGGTT	1,00			visinin-like 1
6042	AAAGGAATAAT	1,00	0,28	Hs.172089	Homo sapiens mRNA; cDNA DKFZp586l2022 (from clone DK
6043	ATTTTAACAAA	1,00	0,28	Hs.155218	E1B-55kDa-associated protein 5
	AACCAGGTGTG	1,00			polymerase (RNA) mitochondrial (DNA directed)
6045	TTTGTGTCAAA	1,00	0,28	Hs.15093	hypothetical protein
6046	GGTCCCGTTCC	1,00			ESTs, Weakly similar to S57447 HPBRII- 7 protein - hu
6047	ACAAAAGACAA	1,00	0,28	Hs.103391	Human insulin-like growth factor binding protein 5 (
6048	CCTTGGTTTTG	3,00	0,76	Hs.75875	ubiquitin-conjugating enzyme E2 variant
6049	TTCTCCCAAAT	3,00	0,76	Hs.75617	collagen, type IV, alpha 2
	TGGTAGTTACC	3,00		Hs.66881	Homo sapiens mRNA; cDNA

					DKFZp434A1518 (from clone DK
	TTTGTTAAAAC	3,00			hypothetical protein
6052	CTTCGAAACTC	4,00	0,96	Hs.51299	NADH dehydrogenase (ubiquinone) flavoprotein 2 (24kD
6053	GCAGTCGCTTG	5,00	1,15	Hs.100002	HSPC162 protein
6054	GCTGAAGGAAA	2,00	0,55	Hs.75569	v-rel avian reticuloendotheliosis viral oncogene hom
6055	GCCTCCTGAGT	2,00	0,55	Hs.108689	sterol regulatory element binding transcription fact
6056	TGAGGCCAGGC	4,00	0,97	Hs.79162	structure specific recognition protein 1
6057	GTCTGGGGCTT	16,00	2,93		transgelin 2
6058	TTGGGGTTTCC	83,00	12,69		ferritin, heavy polypeptide 1
6059	TCAGCCTTCTG	6,00	1,34	Hs.179986	
6060	ATCGCTTTCTA	14,00	2,65	Hs.177486	amyloid beta (A4) precursor protein (protease nexin-
6061	CTTTTGTTTGG	2,00	0,56	Hs.5094	ring finger protein 10
6062	CAAATGCAAAG	2,00	0,56	Hs.11817	nudix (nucleoside diphosphate linked moiety X)-type
6063	AGCACTGCAGC	2,00	0,56	Hs.111039	N-myristoyltransferase 1
6064	TAGGAAACACC	1,00		Hs.8765	RNA helicase-related protein
6065	GACGTTCACTG	1,00	0,28	Hs.75888	phosphogluconate dehydrogenase
6066	TATGCTGTTAT	1,00		Hs.5663	ESTs
6067	TTAGTCAGGCT	1,00			sarcoma amplified sequence
6068	GCCCATTTTAT	1,00	0,28		carbon catabolite repressor protein (CCR4)-associati
6069	GAACGCTGAAG	1,00	0,28	Hs.155983	KIAA0677 gene product
6070	TATCCTAGGGT	1,00			Homo sapiens cDNA FLJ10281 fis, clone HEMBB1001289
6071	TCAACAGCAGG	1,00	0,28	Hs.11556	ESTs
6072	TCCATCAAGAA	6,00	1,36	Hs.79387	proteasome (prosome, macropain) 26S subunit, ATPase,
6073	TAGCCGCTGAG	4,00	0,99	Hs.75932	N-ethylmaleimide-sensitive factor attachment protein
6074	GGCCCTCTGAG	5,00	1,18	Hs.161362	protein (peptidyl-prolyl cis/trans isomerase) NIMA-i
6075	GGTCCAGTGTT	14,00	2,70	Hs.181013	phosphoglycerate mutase 1 (brain)
6076	GGTGGAGCAGA	2,00	0,56	Hs.6846	Homo sapiens mRNA; cDNA DKFZp761J1410 (from clone DK
6077	GCGGCGAG	2,00	0,56	Hs.5662	guanine nucleotide binding protein (G protein), beta
6078	TGTGGGAAATC	2,00	0,56	Hs.251754	secretory leukocyte protease inhibitor (antileukopro
6079	AGTTCCACCAG	2,00	0.56	Hs.182626	chromosome 22 open reading frame 5
	GATGTTAATTG	2,00			ubiquitin specific protease 11
	AACTCTTGAAG	7,00			eukaryotic translation initiation factor 3, subunit
6082	GAAGATGTGGG	11,00	2,22	Hs.250911	Homo sapiens clone 23967 unknown mRNA, partial cds
6083	AACAGAAGCAA	6,00	1.38	Hs.74649	cytochrome c oxidase subunit VIc
		لتتيني			

6094	GTGACAGACAT	2.00	0.70	Uo 75117	interleukin enhancer binding factor 2,
		3,00			45kD
6085	GCTCAGCTGGA	10,00	2,07	Hs.223241 	eukaryotic translation elongation factor 1 delta (gu
6086	TGCTTTGGGAT	7,00	1,56	Hs.84344	CGI-135 protein
6087	GGCTCCTCGAG	4,00	1,00	Hs.179600	TAP binding protein (tapasin)
6088	GCTCCGAGCGT	5,00		Hs.80617	ribosomal protein S16
6089	CCTAGCTGGAT	48,00	8,01	Hs.182937	peptidylprolyl isomerase A (cyclophilin A)
6090	GTGGACCCCAA	4,00	1,01	Hs.74562	siah binding protein 1; FBP interacting
					repressor; p
	GCTTTACTTTG	3,00	0,80	Hs.8966	integral membrane protein 1
	ACCTTCCTAGT	3,00		Hs.6454	chromosome 19 open reading frame 3
6093	AGGAAAAGATG	2,00	0,57	Hs.82520	Human 1.1 kb mRNA upregulated in
					retinoic acid treat
	GTAGGAGCTGG	2,00		Hs.81728	unc119 (C.elegans) homolog
6095	ATTGACCGCTG	2,00	0,57	Hs.177766	ADP-ribosyltransferase (NAD+; poly
					(ADP-ribose) poly
	GAGGACTTGCG	1,00		Hs.8768	hypothetical protein FLJ10849
	GCATTTTGTGA	1,00			TOLLIP protein
	GGCTGGGTTTT	1,00		Hs.74870	H2.0 (Drosophila)-like homeo box 1
	CGTGAACAATT	1,00		Hs.7045	GL004 protein
	TTCCCAAAGGC	1,00			ADP-ribosylation factor related protein 1
6101	TTCACCAGGGC	1,00	0,29	Hs.63243	Homo sapiens cDNA FLJ10041 fis, clone HEMBA1001022
	GTACTCTACTT	1,00	0,29	Hs.52184	hypothetical protein FLJ20618
	CTGGTGGTGCC	1,00	0,29		Gem-interacting protein
	TTCCCTGGGAA	1,00	0,29	Hs.35096	KIAA1538 protein
6105	CTGGGGGAGGG	1,00	0,29	Hs.274122	erythrocyte membrane protein band 4.9 (dematin)
6106	GCTGTACAAAG	1,00	0,29	Hs.250911	Homo sapiens clone 23967 unknown mRNA, partial cds
6107	TGACCTATTTC	1,00	0,29		KIAA0447 gene product
6108	ATCAGTGTGAA	1,00			calponin 3, acidic
6109	AGGAAAGCCAG	1,00	0,29	Hs.19012	Rab9 effector p40
6110	CCGAAGTCGAG	1,00		Hs.169872	
	GAGAAAGAGGC	1,00	0,29	Hs.16420	Npw38-binding protein NpwBP
6112	GCACCTATTGA	1,00		Hs.16178	apoptosis antagonizing transcription factor
6113	TCTTCGTCCTG	1,00	0,29		myelin protein zero-like 1
	CCACCGCACTT	1,00			RAB7, member RAS oncogene family-
0445	0440705151				like 1
6115	GAACTGGAGAA	1,00	0,29	Hs.109150	SH3-domain binding protein 5 (BTK-associated)
	TTGGGCACTAG	1,00	0,29	Hs.102402	Mad4 homolog
	GTTGACTTACA	1,00			hypothetical protein 23851
	AACTTGCCCAT	8,00			ribosomal protein L27
	GTGTTGCACAA	20,00			ribosomal protein S13
6120	TGGAGAGCAAC	4,00	1,01	Hs.4113	S-adenosylhomocysteine hydrolase-like
					· · · · · · · · · · · · · · · · · · ·

10	<u> </u>	T				
by monoclona   b   b   b   b   b   b   b   b   b			4,00			(C/EBP), delta
6123 GAGGGCGTGGG         2,00         0,58 Hs.76368         BCL2-antagonist of cell death           6124 GTGGGGTGACA         2,00         0,58 Hs.183706 adducin 1 (alpha)           6125 TTTTACTCACA         2,00         0,58 Hs.183706 adducin 1 (alpha)           6126 CAATAAACTGA         11,00         2,29 Hs.150580 putative translation initiation factor           6127 ACTGAAGGCGC         3,00         0,81 Hs.92208 a disintegrin and metalloproteinase domain 15 (metar           6128 CTTCTCACCGT         3,00         0,81 Hs.84285 (homologous to yeas           6129 TGGAACTGTGA         3,00         0,81 Hs.279751 sialic acid binding Ig-like lectin 8           6130 AAGAATCTGAA         3,00         0,81 Hs.183435 has a discide by departed	6122	TGACTGGCAGT	4,00	1,02	Hs.119663	
6124 GTGGGGTGACA         2,00         0,58 Hs.250905 hypothetical protein           6125 TTTTACTCACA         2,00         0,58 Hs.183706 adducin 1 (alpha)           6126 CAATAAACTGA         11,00         2,29 Hs.150580 putative translation initiation factor           6127 ACTGAAGGCGC         3,00         0,81 Hs.92208 and disintegrin and metalloproteinase domain 15 (metar           6128 CTTCTCACCGT         3,00         0,81 Hs.279751 slaic acid binding Ig-like lectin 8           6129 TGGAACTGTGA         3,00         0,81 Hs.279751 slaic acid binding Ig-like lectin 8           6130 AAGAATCTGAA         3,00         0,81 Hs.108876 Vacuolar proton-ATPase, subunit Dr.V-ATPase, subunit Dr	6123	GAGGCGCTGGG	2,00	0,58	Hs.76366	BCL2-antagonist of cell death
6125 ITTTACTCACA         2,00         0.58 Hs.183706 ladducin 1 (alpha)           6126 CAATAAACTGA         11,00         2,29 Hs.150580 putative translation initiation factor           6127 ACTGAAGGCGC         3,00         0.81 Hs.92208 a disintegrin and metalloproteinase domain 15 (metar ubiquitin-conjugating enzyme E2I (homologous to yeas           6128 CTTCTCACCGT         3,00         0.81 Hs.84285 (homologous to yeas           6129 TGGAACTGTGA         3,00         0.81 Hs.279751 slalic acid binding Ig-like lectin 8           6130 AAGAATCTGAA         3,00         0.81 Hs.183435 NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 1           6131 GGAAGGACAGA         4,00         1,03 Hs.106876 Vacuolar proton-ATPase, subunit D; V-ATPase, subunit TAPAACCTCAA           6132 GTAAGATTTGA         6,00         1,42 Hs.279939 mitochondrial carrier homolog 1 heterogeneous nuclear ribonucleoproteir D-Ilke           6134 CACCACGGTGT         3,00         0,82 Hs.241471 RNB6           6135 GTAGCAAAAAA         2,00         0,58 Hs.183842 ubiquitin B           6136 GTCCCAAAATG         1,00         0,30 Hs.98614 ribosome binding protein 1 (dog 180kD homolog)           6138 GTCAGACTGTA         1,00         0,30 Hs.93199 ilanosterol synthase (2,3-oxidosqualene-lanosterol cy proteasome (prosome, macropain) 26S subunit, non-ATP independent of the protein 1 displanation and protein 2 displanation and protein 2 displanation and protein 2 displanation 2 displanation 2 displanation 2 displanation 2 displanation 2 disp	6124	GTGGGGTGACA		0,58	Hs.250905	hypothetical protein
6126 CAATAAACTGA         11,00         2,29 Hs. 150580 putative translation initiation factor           6127 ACTGAAGGCGC         3,00         0,81 Hs. 92208 a disintegrin and metaliloproteinase domain 15 (metar           6128 CTTCTCACCGT         3,00         0,81 Hs. 82265 ubiquitin-conjugating enzyme E2I (homologous to yeas           6129 TGGAACTGTGA         3,00         0,81 Hs. 279751 isalia coid binding Ig-like lectin 8           6130 AAGAATCTGAA         3,00         0,81 Hs. 183435 NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 1           6131 GGAAGGACAGA         4,00         1,03 Hs. 106876 Vacuolar proton-ATPase, subunit D; V-ATPase, subunit 10; V-ATPase, subunit 1	6125	TTTTACTCACA				
ACTGAAGGCGC   3,00						
(homologous to yeas   homologous to yeas   homolo	6127	ACTGAAGGCGC				a disintegrin and metalloproteinase
6130 AAGAATCTGAA         3,00         0,81 Hs.183435 NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 1           6131 GGAAGGACAGA         4,00         1,03 Hs.106876 Vacuolar proton-ATPase, subunit D; V-ATPase, subunit Di; V-ATPas		·	3,00			(homologous to yeas
beta subcomplex, 1	6129	TGGAACTGTGA	3,00	0,81	Hs.279751	sialic acid binding Ig-like lectin 8
6131 GGAAGGACAGA 4,00 1,03 Hs.106876 Vacuolar proton-ATPase, subunit D; V-ATPase, subunit Distance in Distance Protein D; V-ATPase, subunit D; V-ATPase, sub	6130	AAGAATCTGAA	3,00	0,81	Hs.183435	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 1
6132         GTAAGATTTGA         6,00         1,42 Hs.279939 mitochondrial carrier homolog 1           6133         TTAAACCTCAA         7,00         1,61 Hs.170311 heterogeneous nuclear ribonucleoproteir D-like           6134         CACCACGGTGT         3,00         0,82 Hs.241471 RNB6           6135         GTAGCAAAAAA         2,00         0,58 Hs.183842 ubiquitin B           6136         GTCCCAAAATG         1,00         0,30 Hs.99908 nuclear receptor coactivator 4           6137         TCGGTTACAAG         1,00         0,30 Hs.98614 ribosome binding protein 1 (dog 180kD homolog)           6138         GTCAGACTGTA         1,00         0,30 Hs.93199 lanosterol synthase (2,3-oxidosqualenelanosterol cy           6139         TAATCTTACT         1,00         0,30 Hs.90744 proteasome (prosome, macropain) 26S subunit, non-ATP           6140         TTAGTCTCAG         1,00         0,30 Hs.82712 fragile X mental retardation, autosomal homolog 1           6141         TCATAGTTCAG         1,00         0,30 Hs.77910 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so           6142         TTTGGAGCATT         1,00         0,30 Hs.7773 ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN           6143         GGAGAGAAAAC         1,00         0,30 Hs.180532 heat shock 90kD protein 1, alpha           6144         GGCCAGTGTTG         1,00         0,30	6131	GGAAGGACAGA	4,00	1,03	Hs.106876	Vacuolar proton-ATPase, subunit D; V-
6133 TTAAACCTCAA         7,00         1,61 Hs. 170311 heterogeneous nuclear ribonucleoprotein D-like           6134 CACCACGGTGT 6135 GTAGCAAAAAA 2,00 0,58 Hs. 183842 ubiquitin B         2,00 0,58 Hs. 183842 ubiquitin B           6136 GTCCCAAAATG 1,00 0,30 Hs. 99908 nuclear receptor coactivator 4         1,00 0,30 Hs. 99908 nuclear receptor coactivator 4 ribosome binding protein 1 (dog 180kD homolog)           6138 GTCAGACTGTA 1,00 0,30 Hs. 93199 lanosterol synthase (2,3-oxidosqualenelanosterol cy proteasome (prosome, macropain) 26S subunit, non-ATP fragile X mental retardation, autosomal homolog 1           6140 TTAGTCTTCAG 1,00 0,30 Hs. 77910 synthase 1 (so         3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so           6142 TTTGGAGCATT 1,00 0,30 Hs. 7773 ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN         ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN           6143 GGAGAGAAAAC 1,00 0,30 Hs. 4094 GGCCAGTGTTG 1,00 0,30 Hs. 4094 Human DNA sequence from clone 167A19 on chromosome 1         Human DNA sequence from clone 167A19 on chromosome 1           6145 CTACTCTTCTA 1,00 0,30 Hs. 169370 FYN oncogene related to SRC, FGR, YES         6148 AGGGGAAAATA 1,00 0,30 Hs. 144477 hypothetical protein PRO2975           6148 AGGGGAAAATA 1,00 0,30 Hs. 104335 Human DNA sequence from clone RP3-402G11 on chromoso         6150 GCCTGCTGGGC 16,00 3,22 Hs. 2706 glutathione peroxidase 4 (phospholipid hydroperoxida           6150 GCCTGCTGGGC 16,00 1,05 Hs. 117582 CGI-43 protein         1,05 Hs. 117582 CGI-43 protein	6132	GTAAGATTTGA	6,00	1,42	Hs.279939	
6135 GTAGCAAAAAA         2,00         0,58 Hs.183842 ubiquitin B           6136 GTCCCAAAATG         1,00         0,30 Hs.99908 nuclear receptor coactivator 4           6137 TCGGTTACAAG         1,00         0,30 Hs.99801 ribosome binding protein 1 (dog 180kD homolog)           6138 GTCAGACTGTA         1,00         0,30 Hs.93199 lanosterol synthase (2,3-oxidosqualene-lanosterol cy           6139 TAATCTTACT         1,00         0,30 Hs.90744 proteasome (prosome, macropain) 26S subunit, non-ATP           6140 TTAGTCTCAG         1,00         0,30 Hs.82712 fragile X mental retardation, autosomal homolog 1           6141 TCATAGTTCAG         1,00         0,30 Hs.77910 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so           6142 TTTGGAGCATT         1,00         0,30 Hs.7773 ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN           6143 GGAGAGAAAAC         1,00         0,30 Hs.70945 ESTs           6144 GGCCAGTGTTG         1,00         0,30 Hs.40094 Human DNA sequence from clone 167A19 on chromosome 1           6145 CTACTCTTCTA         1,00         0,30 Hs.180532 PYN oncogene related to SRC, FGR, YES           6147 TATGTATGTTG         1,00         0,30 Hs.19537 GAP-associated tyrosine phosphoproteir p62 (Sam68)           6149 ATGGCAGAGAC         1,00         0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso           6150 GCCTGCTGGGC         16,00         3,22 Hs.2706 glutathione peroxidase	6133	TTAAACCTCAA				heterogeneous nuclear ribonucleoprotein
6136 GTCCCAAAATG         1,00         0,30 Hs.99908         nuclear receptor coactivator 4           6137 TCGGTTACAAG         1,00         0,30 Hs.98614         ribosome binding protein 1 (dog 180kD homolog)           6138 GTCAGACTGTA         1,00         0,30 Hs.93199         lanosterol synthase (2,3-oxidosqualene-lanosterol cy           6139 TAATCTTACT         1,00         0,30 Hs.90744         proteasome (prosome, macropain) 26S subunit, non-ATP           6140 TTAGTCTCAG         1,00         0,30 Hs.82712         fragile X mental retardation, autosomal homolog 1           6141 TCATAGTTCAG         1,00         0,30 Hs.77910         3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so           6142 TTTGGAGCATT         1,00         0,30 Hs.7793         ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN           6143 GGAGAGAAAAC         1,00         0,30 Hs.70945         ESTs           6144 GGCCAGTGTTG         1,00         0,30 Hs.40094         Human DNA sequence from clone 167A19 on chromosome 1           6145 CTACTCTTCTA         1,00         0,30 Hs.180532 heat shock 90kD protein 1, alpha           6146 TTTTAACAAAA         1,00         0,30 Hs.19370 FYN oncogene related to SRC, FGR, YES           6147 TATGTATGTTG         1,00         0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso           6149 ATGGCAGAGAC         1,00         0,30 Hs.104335 Human	6134	CACCACGGTGT	3,00	0,82	Hs.241471	RNB6
6136 GTCCCAAAATG 1,00 0,30 Hs.99908 nuclear receptor coactivator 4 6137 TCGGTTACAAG 1,00 0,30 Hs.98614 ribosome binding protein 1 (dog 180kD homolog) 6138 GTCAGACTGTA 1,00 0,30 Hs.93199 lanosterol synthase (2,3-oxidosqualene-lanosterol cy proteasome (prosome, macropain) 26S subunit, non-ATP 6140 TTAGTCTTCAG 1,00 0,30 Hs.82712 fragile X mental retardation, autosomal homolog 1 6141 TCATAGTTCAG 1,00 0,30 Hs.77910 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so 6142 TTTGGAGCATT 1,00 0,30 Hs.7793 ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN 6143 GGAGAGAAAAC 1,00 0,30 Hs.70945 ESTs 6144 GGCCAGTGTTG 1,00 0,30 Hs.40094 Human DNA sequence from clone 167A19 on chromosome 1 6145 CTACTCTTCTA 1,00 0,30 Hs.180532 heat shock 90kD protein 1, alpha 6146 TTTTAACAAAA 1,00 0,30 Hs.169370 FYN oncogene related to SRC, FGR, YES 6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso 6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida 66151 GTTTCTCCCT 4,00 1,05 Hs.117582 CGI-43 protein	6135	GTAGCAAAAA	2,00	0,58	Hs.183842	ubiquitin B
6137 TCGGTTACAAG         1,00         0,30 Hs.98614         ribosome binding protein 1 (dog 180kD homolog)           6138 GTCAGACTGTA         1,00         0,30 Hs.93199         lanosterol synthase (2,3-oxidosqualene-lanosterol cy           6139 TAATCTTACT         1,00         0,30 Hs.90744         proteasome (prosome, macropain) 26S subunit, non-ATP fragile X mental retardation, autosomal homolog 1           6140 TTAGTCTCAG         1,00         0,30 Hs.77910         3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so           6142 TTTGGAGCATT         1,00         0,30 Hs.7791         3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so           6143 GGAGAGAAAC         1,00         0,30 Hs.7793         ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN           6144 GCCAGTGTTG         1,00         0,30 Hs.70945         ESTs           6145 CTACTCTTCTA         1,00         0,30 Hs.180532 heat shock 90kD protein 1, alpha           6146 TTTAACAAAA         1,00         0,30 Hs.169370 FYN oncogene related to SRC, FGR, YES           6147 TATGTATGTTG         1,00         0,30 Hs.19537 GAP-associated tyrosine phosphoprotein p62 (Sam68)           6149 ATGGCAGAGAC         1,00         0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso           6150 GCCTGCTGGGC         16,00         3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida           6151 GTTTCTCCCT         4,00 <td>6136</td> <td>GTCCCAAAATG</td> <td>1,00</td> <td></td> <td></td> <td></td>	6136	GTCCCAAAATG	1,00			
lanosterol cy 6139 TAATCTTACT 1,00 0,30 Hs.90744 proteasome (prosome, macropain) 26S subunit, non-ATP 6140 TTAGTCTCAG 1,00 0,30 Hs.82712 fragile X mental retardation, autosomal homolog 1 6141 TCATAGTTCAG 1,00 0,30 Hs.77910 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (so 6142 TTTGGAGCATT 1,00 0,30 Hs.7773 ESTs, Weakly similar to A4P_HUMAN INTESTINAL MEMBRAN 6143 GGAGAGAAAC 1,00 0,30 Hs.70945 ESTs 6144 GGCCAGTGTTG 1,00 0,30 Hs.40094 Human DNA sequence from clone 167A19 on chromosome 1 6145 CTACTCTTCTA 1,00 0,30 Hs.180532 heat shock 90kD protein 1, alpha 6146 TTTAACAAAA 1,00 0,30 Hs.169370 FYN oncogene related to SRC, FGR, YES 6147 TATGTATGTTG 1,00 0,30 Hs.144477 hypothetical protein PRO2975 6148 AGGGGAAAATA 1,00 0,30 Hs.119537 GAP-associated tyrosine phosphoprotein p62 (Sam68) 6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso 6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida 6151 GTTTCTTCCCT 4,00 1,05 Hs.117582 CGI-43 protein	6137	TCGGTTACAAG				ribosome binding protein 1 (dog 180kD
Subunit, non-ATP			1,00			
homolog 1			1,00	0,30	Hs.90744	
Synthase 1 (so   6142 TTTGGAGCATT   1,00   0,30 Hs.7773   ESTs, Weakly similar to A4P_HUMAN   INTESTINAL MEMBRAN   6143 GGAGAGAAAC   1,00   0,30 Hs.70945   ESTs	6140	TTAGTCTTCAG	1,00	0,30	Hs.82712	homolog 1
INTESTINAL MEMBRAN   6143 GGAGAGAAAC   1,00   0,30   Hs.70945   ESTs   6144   GGCCAGTGTTG   1,00   0,30   Hs.40094   Human DNA sequence from clone   167A19 on chromosome 1   6145 CTACTCTTCTA   1,00   0,30   Hs.180532   heat shock 90kD protein 1, alpha   6146   TTTTAACAAAA   1,00   0,30   Hs.169370   FYN oncogene related to SRC, FGR, YES   6147 TATGTATGTTG   1,00   0,30   Hs.144477   hypothetical protein PRO2975   6148   AGGGGAAAATA   1,00   0,30   Hs.119537   GAP-associated tyrosine phosphoprotein   p62 (Sam68)   6149   ATGGCAGAGAC   1,00   0,30   Hs.104335   Human DNA sequence from clone RP3-402G11 on chromoso   6150   GCCTGCTGGGC   16,00   3,22   Hs.2706   glutathione peroxidase 4 (phospholipid hydroperoxida   6151   GTTTCTTCCCT   4,00   1,05   Hs.117582   CGI-43 protein	6141	TCATAGTTCAG	1,00	0,30	Hs.77910	synthase 1 (so
6144 GGCCAGTGTTG 1,00 0,30 Hs.40094 Human DNA sequence from clone 167A19 on chromosome 1 6145 CTACTCTTCTA 1,00 0,30 Hs.180532 heat shock 90kD protein 1, alpha 6146 TTTTAACAAAA 1,00 0,30 Hs.169370 FYN oncogene related to SRC, FGR, YES 6147 TATGTATGTTG 1,00 0,30 Hs.144477 hypothetical protein PRO2975 6148 AGGGGAAAATA 1,00 0,30 Hs.119537 GAP-associated tyrosine phosphoprotein p62 (Sam68) 6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso 6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida	6142	TTTGGAGCATT	1,00	0,30	Hs.7773	
167A19 on chromosome 1 6145 CTACTCTTCTA 1,00 0,30 Hs.180532 heat shock 90kD protein 1, alpha 6146 TTTTAACAAAA 1,00 0,30 Hs.169370 FYN oncogene related to SRC, FGR, YES 6147 TATGTATGTTG 1,00 0,30 Hs.144477 hypothetical protein PRO2975 6148 AGGGGAAAATA 1,00 0,30 Hs.119537 GAP-associated tyrosine phosphoprotein p62 (Sam68) 6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3- 402G11 on chromoso 6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida 6151 GTTTCTTCCCT 4,00 1,05 Hs.117582 CGI-43 protein			1,00	0,30	Hs.70945	
6146 TTTTAACAAAA 1,00 0,30 Hs.169370 FYN oncogene related to SRC, FGR, YES 6147 TATGTATGTTG 1,00 0,30 Hs.144477 hypothetical protein PRO2975 6148 AGGGGAAAATA 1,00 0,30 Hs.119537 GAP-associated tyrosine phosphoprotein p62 (Sam68) 6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso 6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida 6151 GTTTCTTCCCT 4,00 1,05 Hs.117582 CGI-43 protein	6144	GGCCAGTGTTG	1,00	0,30	Hs.40094	
YES  6147 TATGTATGTTG 1,00 0,30 Hs.144477 hypothetical protein PRO2975  6148 AGGGGAAAATA 1,00 0,30 Hs.119537 GAP-associated tyrosine phosphoprotein p62 (Sam68)  6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso  6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida  6151 GTTTCTTCCCT 4,00 1,05 Hs.117582 CGI-43 protein						
6148 AGGGAAAATA 1,00 0,30 Hs.119537 GAP-associated tyrosine phosphoprotein p62 (Sam68) 6149 ATGGCAGAGAC 1,00 0,30 Hs.104335 Human DNA sequence from clone RP3-402G11 on chromoso 6150 GCCTGCTGGGC 16,00 3,22 Hs.2706 glutathione peroxidase 4 (phospholipid hydroperoxida 6151 GTTTCTTCCCT 4,00 1,05 Hs.117582 CGI-43 protein	٠					YES
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6151 GTTTCTTCCCT 4,00 1,05 Hs.117582 CGI-43 protein			16,00	3,22	Hs.2706	
	6151	GTTTCTTCCCT	4,00	1,05	Hs.117582	
	6152	CCGGACCTGTG				

	GCTTTCATTGG	5,00			Homo sapiens clone 23967 unknown mRNA, partial cds
6154	TTGGAGATCTC	19,00		Hs.50098	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,
6155	ACTGCTGAACC	3,00	0,83	Hs.200600	secretory carrier membrane protein 3
6156	CACTTGCCCTA	11,00		Hs.15977	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 9
6157	GCCAAGATGCC	4,00	1,06	Hs.83135	p53-responsive gene 6
	CCTGCCCCCCT	3,00		Hs.861	mitogen-activated protein kinase 3
	ACACTTCTTTC	2,00		Hs.83381	guanine nucleotide binding protein 11
	GCTAGGTCTGG	2,00		Hs.75354	GCN1 (general control of amino-acid synthesis 1, yea
6161	TCCATCCCTTG	2,00	0,60	Hs.7527	small fragment nuclease
6162	AGGCCTGGCTA	2,00	0,60	Hs.5011	RNA binding motif protein 9
6163	GAGAGCCTGCC	2,00	0,60	Hs.31305	transducin-like enhancer of split 3, homolog of Dros
6164	GTTTAAAAAGA	1,00	0,30	Hs.90005	superiorcervical ganglia, neural specific 10
6165	CCAAGGACTCT	1,00	0,30	Hs.79058	suppressor of Ty (S.cerevisiae) 4 homolog 1
6166	GAAAAGTTGCC	1,00	0,30	Hs.33122	KIAA1160 protein
6167	AATTATGACTT	1,00		Hs.2853	poly(rC)-binding protein 1
6168	TCAGTGCTCTC	1,00	0,30	Hs.284741	Human clone 23745 mRNA, complete cds
6169	CCTGTCTGCAC	1,00	0,30	Hs.25338	ESTs
6170	CGAATAAAATG	1,00		Hs.20815	erythroblast macrophage protein
6171	TTTGGACAATA	1,00	0,30	Hs.173685	Human DNA sequence from clone 30M3 on chromosome 6p2
6172	GACGACTGACC	1,00	0,30	Hs.155530	interferon, gamma-inducible protein 16
6173	CCCTATCACAA	1,00			CATX-8 protein
6174	GTGAATAAACA	1,00			Homo sapiens cDNA FLJ20678 fis, clone KAIA4163
6175	GTGGGACCATT	1,00		*	protein (peptidyl-prolyl cis/trans isomerase) NIMA-i
	CTCAGGAAATA	7,00			ubiquinol-cytochrome c reductase complex (7.2 kD)
6177	ACCAAGGAGGA	5,00	1,27	Hs.283547	HSVI binding protein
6178	TCTCTACCCAC	8,00	1,87	Hs.64797	amyloid beta (A4) precursor-like protein 2
6179	CAGTCTCTCAA	10,00	2,25	Hs.76230	ribosomal protein S10
6180	CCTTGAGTACA	4,00	1,09	Hs.155247	aldolase C, fructose-bisphosphate
	TGCAGACCCAT	2,00		Hs.5437	Tax1 (human T-cell leukemia virus type I) binding pr
	GGGCCAAAACC	1,00		Hs.9142	ESTs, Weakly similar to FLDED-1 [H.sapiens]
6183	GTCCTTCAGAA	1,00	. 0,31	Hs.82712	fragile X mental retardation, autosomal homolog 1
6184	CAGTCTGGGAG	1,00	0,31	Hs.285115	interleukin 13 receptor, alpha 1
	CGCCGCTTCTT	1,00			bromodomain adjacent to zinc finger

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6220         CCCCCGTGAAG         5,00         1,36         Hs.182018 interleukin-1 receptor-associated kina feezt TCTTCCAGGAG         6,00         1,58         Hs.74267 inbosomal protein L15           6221         TCTTCCAGGAG         6,00         1,58         Hs.74267 inbosomal protein L15           6222         CATTAAATTCA         2,00         0,64         Hs.31053 cytoskeleton-associated protein 1           6224         GGGGCTGTATT         2,00         0,64         Hs.146393 [KlAA0025 gene product; MMS-induct gene           6224         GGGGCTGTATT         2,00         0,64         Hs.146393 [KlAA0025 gene product; MMS-induct gene           6225         TGGAGGCCAGG         3,00         0,90         Hs.250581 [SW/INSNF related, matrix associated, actin dependent           6226         TGTGGGTGCTG         4,00         1,14 [Hs.194657] cadherin 1, E-cadherin (epithelial)           6227         TGGGAGGATTAA         1,00         0,32 [Hs.90783] Tat-interacting protein (30kD)           6228         GTGTGATGCTG         1,00         0,32 [Hs.807893] prolylcarboxypeptidase (angiotensina cyberlada) [Carl Adole protein actinal protein protein actinal protein act						isoform
1	6220	CCCCCGTGAAG	5.00	1 36	He 182018	
6222 CATTAAATTCA         2,00         0,64 Hs.31053 cytoskeleton-associated protein 1           6223 TTTTAGCAGGA         2,00         0,64 Hs.146393 kt/AA0025 gene product; MMS-induct gene           6224 GGGCTGTATT         2,00         0,64 Hs.1103 transforming growth factor, beta 1           6225 TGGAAGCCAGG         3,00         0,90 Hs.250581 SWW/SNF related, matrix associated, actin dependent           6226 TGTGGGTGCTG         4,00         1,14 Hs.194657 cadherin 1, E-cadherin (epithelial)           6227 GGGAGGATTAA         1,00         0,32 Hs.90753         Tat-interacting protein (30kD)           6228 GTGTGATCTC         1,00         0,32 Hs.75693 prolylcarboxypeptidase (angiotensina C)           6230 ACATCATACTG         1,00         0,32 Hs.260666 Homo sapiens chromosome 19, cosn R32184           6231 TGATGTTTGCA         1,00         0,32 Hs.280666 Homo sapiens chromosome 19, cosn R32184           6233 GTTGGATAGGG         1,00         0,32 Hs.225851 aspartly aminopeptidase           6234 CACTCTATCCG         1,00         0,32 Hs.27184 growth factor, erv1 (S. cerevisiae)-like (augmenter           6234 CACTCTATCGG         1,00         0,32 Hs.27184 growth factor pathway inhibitor (lipoprotein-associa           6235 AATTCTGTAAA         1,00         0,32 Hs.170279 fissue factor pathway inhibitor (lipoprotein-associa)           6236 CTGAAAACCAC         1,00         0,32 Hs.168350 kliAA0554 protein						1
6223         TTTTAGCAGGA         2,00         0,64 Hs.146393         KIAA0025 gene product; MMS-inducion gene           6224         GGGGCTGTATT         2,00         0,64 Hs.1103         transforming growth factor, beta 1           6225         TGGAGGCCAGG         3,00         0,90 Hs.250581         SWI/SNF related, matrix associated, actin dependent           6226         TGTGGGTGCTG         4,00         1,14 Hs.194657         cadherin 1, E-cadherin (epithelial)           6227         GGGAGGATTAA         1,00         0,32 Hs.90753         Tat-interacting protein (30kD)           6228         GTGTGATGCTG         1,00         0,32 Hs.89519         KIAA1046 protein           6229         AATGTAATCTG         1,00         0,32 Hs.61790         ESTs, Weakly similar to T01239 hypothetical protein           6231         TGATGTTTGCA         1,00         0,32 Hs.280666 Homo sapiens chromosome 19, cosn R32184           6233         GTTGGATAGGG         1,00         0,32 Hs.27184         growth factor, erv1 (S. cerevisiae)-like (augmenter           6234         CACTCTATCCG         1,00         0,32 Hs.281574         Human DNA sequence from clone 316G12 on chromosome 1           6236         CTGAAAACCAC         1,00         0,32 Hs.16830         KiAA054 protein           6237         TGCTGAGGAG         1,00 <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td></td<>						
gene   gene   transforming growth factor, beta 1						
6224 GGGCTGTATT         2,00         0,64 Hs.1103         transforming growth factor, beta 1           6225 TGGAGGCCAGG         3,00         0,90 Hs.250581         SW//SNF related, matrix associated, actin dependent           6226 TGTGGGTGCTG         4,00         1,14 Hs.194657         Cadherin 1, E-cadherin (epithelial)           6227 GGGAGGATTAA         1,00         0,32 Hs.90753         Tat-interacting protein (30kD)           6228 GTGTGATGCTG         1,00         0,32 Hs.89519         KIAA1046 protein           6229 AATGTGATTTC         1,00         0,32 Hs.61790         ESTs, Weakly similar to T01239 hypothetical protein           6230 ACATCATACTG         1,00         0,32 Hs.4552         ubiquilin 2           6231 TGATGTTTGCA         1,00         0,32 Hs.280666         Homo sapiens chromosome 19, cosn R32184           6232 CTTCTGTTTT         1,00         0,32 Hs.27184         growth factor, erv1 (S. cerevisiae)-like (augmenter           6233 GTTGGATAGGG         1,00         0,32 Hs.27184         growth factor pathway inhibitor           6234 CACTCTATCCG         1,00         0,32 Hs.265551         aspartyl aminopeptidase           6235 AATTCTGTAAA         1,00         0,32 Hs.168350 KlAA0554 protein           6236 CTGAAAACCAC         1,00         0,32 Hs.168350 KlAA0554 protein           6237 TGCTGAGGAAG         1,00<	6223	TTTTAGCAGGA	2,00	0,64	Hs.146393	
6225 TGGAGGCCAGG         3,00         0,90 Hs.250581 SWI/SNF related, matrix associated, actin dependent           6226 TGTGGGTGCTG         4,00         1,14 Hs.194657 cadherin 1, E-cadherin (epithelial)           6227 GGGAGGATTAA         1,00         0,32 Hs.90753 Tat-interacting protein (30kD)           6228 GTGTGATGCTG         1,00         0,32 Hs.90753 Tat-interacting protein (30kD)           6229 AATGTGATTTC         1,00         0,32 Hs.75693 Protein           6230 ACATCATACTG         1,00         0,32 Hs.61790 Protein prolycarboxypeptidase (angiotensina C)           6231 TGATGTTTGCA         1,00         0,32 Hs.4552 Protein	6224	GGGGCTGTATT	2 00	0.64	Hs 1103	
actin dependent   actin dependent   actin dependent   G226 TGTGGGTGCTG   4,00   1,14   Hs. 194657   cacherin 1, E-cadherin (epithelial)   6227 GGGAGGATTAA   1,00   0,32   Hs. 90753   Tat-interacting protein (30kD)   6228 GTGTGATGCTG   1,00   0,32   Hs. 95519   KIAA1046 protein   6229   AATGTGATTTC   1,00   0,32   Hs. 75693   prolylcarboxypeptidase (angiotensina C)   6230   ACATCATACTG   1,00   0,32   Hs. 61790   ESTs, Weakly similar to T01239   hypothetical protein   6231 TGATGTTTGCA   1,00   0,32   Hs. 4552   ubiquilin 2   6232   CTTCTGTTTT   1,00   0,32   Hs. 28066   Homo sapiens chromosome 19, cosn   R32184   growth factor, erv1 (S. cerevisiae)-like (augmenter   6234   CACTCTATCCG   1,00   0,32   Hs. 28155   aspartyl aminopeptidase   6235   AATTCTGTAAA   1,00   0,32   Hs. 241575   Human DNA sequence from clone   316G12 on chromosome 1   6236   CTGAAAACCAC   1,00   0,32   Hs. 170279   tissue factor pathway inhibitor (lipoprotein-associa   6237 TGCTGAGGAAG   1,00   0,32   Hs. 157145   tetracycline transporter-like protein   6239 TACATATGGAA   1,00   0,32   Hs. 157145   tetracycline transporter-like protein   6239 TACATATGGAA   1,00   0,32   Hs. 157145   tetracycline transporter-like protein   6240 TCTGCTTTTGA   1,00   0,32   Hs. 144504   hypothetical protein FLJ10624   6241   ACAGCCAAGAG   1,00   0,32   Hs. 144504   hypothetical protein FLJ10624   6242 GGCTTGCTGAC   1,00   0,32   Hs. 19403   hexosaminidase A (alpha polypeptide   6245 CAGCCTTGGAC   3,00   0,91   Hs. 65648   RNA binding motif protein 8   6246   ACAGGGTGACC   1,00   2,48   Hs. 174050   endothelial differentiation-related fact   6247 TCCCTGTACAT   5,00   1,40   Hs. 89563   nuclear cap binding protein subunit 1, 80kD   6248   ACTCTGCCAAG   2,00   0,65   Hs. 241543   DKF2F86F1524 protein   6250 CAAATGAGGAG   1,00   0,33   Hs. 83081   GTP cyclohydrolase I feedback   6250 CAAATGAGGAG   1,00   0,33   Hs. 83081   GTP cyclohydrolase I feedback   6250 CAAATGAGGAG   1,00   0,33   Hs. 83081   GTP cyclohydrolase I feedback   6250 CAAATGAGGAG   1,00   0,30			3.00			
6227         GGGAGGATTAA         1,00         0,32 Hs.90753         Tat-interacting protein (30kD)           6228         GTGTGATGCTG         1,00         0,32 Hs.89519         KIAA1046 protein           6229         AATGTGATTC         1,00         0,32 Hs.75693         prolylcarboxypeptidase (angiotensina C)           6230         ACATCATACTG         1,00         0,32 Hs.61790         ESTs, Weakly similar to T01239 hypothetical protein           6231         TGATGTTTGCA         1,00         0,32 Hs.280666         Homo sapiens chromosome 19, cosm R32184           6232         CTTCTGTTTT         1,00         0,32 Hs.27184         growth factor, erv1 (S. cerevisiae)-like (augmenter Growth factor) (S. cerevisiae)-like (Growth factor)						actin dependent
6228 GTGTGATGCTG         1,00         0,32 Hs.89519         KIAA1046 protein           6229 AATGTGATTC         1,00         0,32 Hs.75693         prolycarboxypeptidase (angiotensina C)           6230 ACATCATACTG         1,00         0,32 Hs.61790         ESTs, Weakly similar to T01239 hypothetical protein           6231 TGATGTTTGCA         1,00         0,32 Hs.4552         ubiquilin 2           6232 CTTCTGTTTT         1,00         0,32 Hs.280666         Homo sapiens chromosome 19, cosm R32184           6233 GTTGGATAGGG         1,00         0,32 Hs.27184         growth factor, ev1 (S. cerevisiae)-like (augmenter           6234 CACTCTATCCG         1,00         0,32 Hs.241575         Human DNA sequence from clone 316G12 on chromosome 1           6236 CTGAAAACCAC         1,00         0,32 Hs.170279         tissue factor pathway inhibitor (lipoprotein-associa           6237 TGCTGAGGAAG         1,00         0,32 Hs.168350 KlAA0554 protein           6238 GGTACACTGCG         1,00         0,32 Hs.155499 golgi-specific brefeldin A resistance factor 1           6239 TACATATGGAA         1,00         0,32 Hs.144504 hypothetical protein FLJ10624           6240 TCTGCTTTTGA         1,00         0,32 Hs.144504 hypothetical protein FLJ10624           6241 ACAGCCAAGAG         1,00         0,32 Hs.13699         decay accelerating factor for complen (CD55, Crom           <						
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hypothetical protein	6229	AATGTGATTTC	1,00	0,32	Hs.75693	I i i i i i i i i i i i i i i i i i i i
6231 TGATGTTTGCA         1,00         0,32 Hs.4552         ubiquilin 2           6232 CTTCTGTTTTT         1,00         0,32 Hs.280666 Homo sapiens chromosome 19, cosm R32184           6233 GTTGGATAGGG         1,00         0,32 Hs.27184         growth factor, erv1 (S. cerevisiae)-like (augmenter)           6234 CACTCTATCCG         1,00         0,32 Hs.258551         aspartyl aminopeptidase           6235 AATTCTGTAAA         1,00         0,32 Hs.241575         Human DNA sequence from clone 316G12 on chromosome 1           6236 CTGAAAACCAC         1,00         0,32 Hs.170279         tissue factor pathway inhibitor (lipoprotein-associa           6237 TGCTGAGGAAG         1,00         0,32 Hs.168350 KIAA0554 protein           6238 GGTACACTGCG         1,00         0,32 Hs.157145 tetracycline transporter-like protein           6239 TACATATGGAA         1,00         0,32 Hs.144504 hypothetical protein FLJ10624           6240 TCTGCTTTTGA         1,00         0,32 Hs.144504 hypothetical protein FLJ10624           6241 ACAGCCAAGAG         1,00         0,32 Hs.114422 nudix (nucleoside diphosphate linked moiety X)-type           6242 GGCTTGCTGAC         1,00         0,32 Hs.119403 hexosaminidase A (alpha polypeptide McCD55, Crom           6243 GATTACCTGTG         1,00         0,32 Hs.119403 hexosaminidase A (alpha polypeptide Dratein 1           6244 TCCGTGGTGAC         1,00         <	6230	ACATCATACTG	1,00	0,32	Hs.61790	
6232 CTTCTGTTTTT         1,00         0,32 Hs.280666 Homo sapiens chromosome 19, cosm R32184           6233 GTTGGATAGGG         1,00         0,32 Hs.27184 growth factor, erv1 (S. cerevisiae)-like (augmenter (augmenter (augmenter) (au			4 00			
R32184   G233   GTTGGATAGGG   1,00   0,32   Hs.27184   growth factor, erv1 (S. cerevisiae)-like (augmenter   G234   CACTCTATCCG   1,00   0,32   Hs.258551   aspartyl aminopeptidase   G235   AATTCTGTAAA   1,00   0,32   Hs.241575   Human DNA sequence from clone   316G12 on chromosome 1   G236   CTGAAAACCAC   1,00   0,32   Hs.170279   tissue factor pathway inhibitor (lipoprotein-associa   G237   TGCTGAGGAAG   1,00   0,32   Hs.168350   KIAA0554 protein   G238   GGTACACTGCG   1,00   0,32   Hs.157145   tetracycline transporter-like protein   G239   TACATATGGAA   1,00   0,32   Hs.155499   golgi-specific brefeldin A resistance   factor 1   G241   ACAGCCAAGAG   1,00   0,32   Hs.144504   hypothetical protein   FLJ10624   nudix (nucleoside diphosphate linked moiety X)-type   decay accelerating factor for complen   (CD55, Crom   G243   GATTACCTGTG   1,00   0,32   Hs.119403   hexosaminidase A (alpha polypeptide   G244   TCCGTGGTTGG   7,00   1,81   Hs.79516   brain acid-soluble protein 1   G245   CAGCCTTGGAC   3,00   0,91   Hs.65648   RNA binding motif protein 8   G246   ACAGGGTGACC   10,00   2,48   Hs.174050   endothelial differentiation-related factor   G249   ACTCTGCCAAG   2,00   0,65   Hs.241543   DKFZP586F1524 protein   G250   CAAATGAGGAG   14,00   3,32   Hs.69855   NRAS-related gene   G250   CAAATGAGGAG   14,00   3,32   Hs.69855   NRAS-related gene   G7P cyclohydrolase   feedback   G7P cycl						
(augmenter   6234   CACTCTATCCG   1,00   0,32   Hs.258551   aspartyl aminopeptidase   6235   AATTCTGTAAA   1,00   0,32   Hs.241575   Human DNA sequence from clone 316G12 on chromosome 1   6236   CTGAAAACCAC   1,00   0,32   Hs.170279   tissue factor pathway inhibitor (lipoprotein-associa   6237   TGCTGAGGAAG   1,00   0,32   Hs.168350   KIAAO554   protein   6238   GGTACACTGCG   1,00   0,32   Hs.157145   tetracycline transporter-like protein   6239   TACATATGGAA   1,00   0,32   Hs.155499   golgi-specific brefeldin A resistance factor 1   ACAGCCAAGAG   1,00   0,32   Hs.144504   hypothetical protein FLJ10624   nudix (nucleoside diphosphate linked moiety X)-type   decay accelerating factor for complen (CD55, Crom   6243   GATTACCTGTG   1,00   0,32   Hs.1369   decay accelerating factor for complen (CD55, Crom   6244   TCCGTGGTTGG   7,00   1,81   Hs.79516   brain acid-soluble protein 1   6245   CAGCCTTGGAC   3,00   0,91   Hs.65648   RNA binding motif protein 8   6246   ACAGGGTGACC   1,00   2,48   Hs.174050   endothelial differentiation-related factor for the fa	6232	CTTCTGTTTT	1,00	0,32	Hs.280666	R32184
6234 CACTCTATCCG         1,00         0,32 Hs.258551         aspartyl aminopeptidase           6235 AATTCTGTAAA         1,00         0,32 Hs.241575         Human DNA sequence from clone 316G12 on chromosome 1           6236 CTGAAAACCAC         1,00         0,32 Hs.170279         tissue factor pathway inhibitor (lipoprotein-associa           6237 TGCTGAGGAAG         1,00         0,32 Hs.168350 KlAA0554 protein           6238 GGTACACTGCG         1,00         0,32 Hs.157145 tetracycline transporter-like protein           6239 TACATATGGAA         1,00         0,32 Hs.155499 golgi-specific brefeldin A resistance factor 1           6240 TCTGCTTTTGA         1,00         0,32 Hs.144504 hypothetical protein FLJ10624           6241 ACAGCCAAGAG         1,00         0,32 Hs.14142 nudix (nucleoside diphosphate linked moiety X)-type           6242 GGCTTGCTGAC         1,00         0,32 Hs.119403 hexosaminidase A (alpha polypeptide form of the protein factor for complen (CD55, Crom           6243 GATTACCTGTG         1,00         0,32 Hs.119403 hexosaminidase A (alpha polypeptide form of the protein factor for complen (CD55, Crom           6244 TCCGTGGTTGG         7,00         1,81 Hs.79516 brain acid-soluble protein for the protein factor for complen for the protein factor for complen for the protein factor for complen factor for factor for factor for factor for factor fa	6233	GTTGGATAGGG	1,00	0,32	Hs.27184	
6235         AATTCTGTAAA         1,00         0,32         Hs.241575         Human DNA sequence from clone 316G12 on chromosome 1           6236         CTGAAAACCAC         1,00         0,32         Hs.170279         tissue factor pathway inhibitor (lipoprotein-associa           6237         TGCTGAGGAAG         1,00         0,32         Hs.168350         KIAA0554 protein           6238         GGTACACTGCG         1,00         0,32         Hs.157145         tetracycline transporter-like protein           6239         TACATATGGAA         1,00         0,32         Hs.155499         golgi-specific brefeldin A resistance factor 1           6240         TCTGCTTTTGA         1,00         0,32         Hs.144504         hypothetical protein FLJ10624           6241         ACAGCCAAGAG         1,00         0,32         Hs.14142         nudix (nucleoside diphosphate linked moiety X)-type           6242         GGCTTGCTGAC         1,00         0,32         Hs.119403         hexosaminidase A (alpha polypeptide for complem (CD55, Crom           6243         GATTACCTGTG         1,00         0,32         Hs.119403         hexosaminidase A (alpha polypeptide for complem (CD55, Crom           6244         TCCGTGGTGG         7,00         1,81         Hs.79516         brain acid-soluble protein 1           6245 <td>6234</td> <td>CACTCTATCCG</td> <td>1.00</td> <td>0.32</td> <td>Hs.258551</td> <td></td>	6234	CACTCTATCCG	1.00	0.32	Hs.258551	
6236 CTGAAAACCAC 1,00 0,32 Hs.170279 tissue factor pathway inhibitor (lipoprotein-associa 6237 TGCTGAGGAAG 1,00 0,32 Hs.168350 KIAA0554 protein 6238 GGTACACTGCG 1,00 0,32 Hs.157145 tetracycline transporter-like protein 6239 TACATATGGAA 1,00 0,32 Hs.155499 golgi-specific brefeldin A resistance factor 1 1,00 0,32 Hs.144504 hypothetical protein FLJ10624 6241 ACAGCCAAGAG 1,00 0,32 Hs.144504 hypothetical protein FLJ10624 nudix (nucleoside diphosphate linked moiety X)-type 6242 GGCTTGCTGAC 1,00 0,32 Hs.1369 decay accelerating factor for complem (CD55, Crom 6243 GATTACCTGTG 1,00 0,32 Hs.119403 hexosaminidase A (alpha polypeptide 6244 TCCGTGGTTGG 7,00 1,81 Hs.79516 brain acid-soluble protein 1 6245 CAGCCTTGGAC 3,00 0,91 Hs.65648 RNA binding motif protein 8 6246 ACAGGGTGACC 10,00 2,48 Hs.174050 endothelial differentiation-related factor for CCTGTACAT 5,00 1,40 Hs.89563 nuclear cap binding protein subunit 1,80kD 6248 TTATGGATCTC 5,00 1,40 Hs.5662 guanine nucleotide binding protein (Gprotein), beta 6250 CAAATGAGGAG 14,00 3,32 Hs.69855 NRAS-related gene 6251 TTCCTCGGGCA 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback				0,32	Hs.241575	Human DNA sequence from clone
6237 TGCTGAGGAAG         1,00         0,32 Hs.168350 KIAA0554 protein           6238 GGTACACTGCG         1,00         0,32 Hs.157145 tetracycline transporter-like protein           6239 TACATATGGAA         1,00         0,32 Hs.155499 golgi-specific brefeldin A resistance factor 1           6240 TCTGCTTTTGA         1,00         0,32 Hs.144504 hypothetical protein FLJ10624           6241 ACAGCCAAGAG         1,00         0,32 Hs.14142 nudix (nucleoside diphosphate linked moiety X)-type           6242 GGCTTGCTGAC         1,00         0,32 Hs.1369 decay accelerating factor for complen (CD55, Crom           6243 GATTACCTGTG         1,00         0,32 Hs.119403 hexosaminidase A (alpha polypeptide brain acid-soluble protein 1           6244 TCCGTGGTTGG         7,00         1,81 Hs.79516 brain acid-soluble protein 8           6245 CAGCCTTGGAC         3,00         0,91 Hs.65648 RNA binding motif protein 8           6246 ACAGGGTGACC         10,00         2,48 Hs.174050 endothelial differentiation-related factor           6247 TCCCTGTACAT         5,00         1,40 Hs.89563 nuclear cap binding protein subunit 1, 80kD           6248 TTATGGATCTC         5,00         1,40 Hs.5662 guanine nucleotide binding protein (Gprotein), beta           6249 ACTCTGCCAAG         2,00         0,65 Hs.241543 DKFZP586F1524 protein           6250 CAAATGAGGAG         14,00         3,32 Hs.69855 NRAS-related gene           <	6236	CTGAAAACCAC	1,00	0,32	Hs.170279	tissue factor pathway inhibitor
GCTACACTGCG 1,00 0,32 Hs.157145 tetracycline transporter-like protein 0,32 Hs.155499 golgi-specific brefeldin A resistance factor 1 1,00 0,32 Hs.14504 hypothetical protein FLJ10624 nudix (nucleoside diphosphate linked moiety X)-type decay accelerating factor for complem (CD55, Crom (CD55, Crom 1,00 0,32 Hs.119403 hexosaminidase A (alpha polypeptide 1,00 0,32 Hs.19403 hexosaminidase A (alpha polypeptide 1,00 0,32 Hs.19403 hexosaminidase A (alpha polypeptide 1,00 0,31 Hs.65648 RNA binding motif protein 8 Hs.74050 endothelial differentiation-related factor for CCTGTACAT 5,00 1,40 Hs.89563 nuclear cap binding protein subunit 1,80kD 1,40 Hs.5662 guanine nucleotide binding protein (CD55) CAAATGAGGAG 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback	6227	TGCTGAGGAAG	1.00	0.33	He 169350	
FACATATGGAA  1,00  0,32  Hs. 155499  golgi-specific brefeldin A resistance factor 1  1,00  0,32  Hs. 144504  hypothetical protein FLJ10624  nudix (nucleoside diphosphate linked moiety X)-type  1,00						
factor 1 6240 TCTGCTTTTGA 1,00 0,32 Hs.144504 hypothetical protein FLJ10624 6241 ACAGCCAAGAG 1,00 0,32 Hs.14142 nudix (nucleoside diphosphate linked moiety X)-type 6242 GGCTTGCTGAC 1,00 0,32 Hs.1369 decay accelerating factor for complen (CD55, Crom 6243 GATTACCTGTG 1,00 0,32 Hs.119403 hexosaminidase A (alpha polypeptide 6244 TCCGTGGTTGG 7,00 1,81 Hs.79516 brain acid-soluble protein 1 6245 CAGCCTTGGAC 3,00 0,91 Hs.65648 RNA binding motif protein 8 6246 ACAGGGTGACC 10,00 2,48 Hs.174050 endothelial differentiation-related fact 6247 TCCCTGTACAT 5,00 1,40 Hs.89563 nuclear cap binding protein subunit 1,80kD 6248 TTATGGATCTC 5,00 1,40 Hs.5662 guanine nucleotide binding protein (Gprotein), beta 6249 ACTCTGCCAAG 2,00 0,65 Hs.241543 DKFZP586F1524 protein 6250 CAAATGAGGAG 14,00 3,32 Hs.69855 NRAS-related gene 6251 TTCCTCGGGCA 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback				0,32	Ho 155 140	galai appoific brofoldin A registance
6241 ACAGCCAAGAG 1,00 0,32 Hs.14142 nudix (nucleoside diphosphate linked moiety X)-type 6242 GGCTTGCTGAC 1,00 0,32 Hs.1369 decay accelerating factor for complem (CD55, Crom 6243 GATTACCTGTG 1,00 0,32 Hs.119403 hexosaminidase A (alpha polypeptide form acid-soluble protein 1 6244 TCCGTGGTTGG 7,00 1,81 Hs.79516 brain acid-soluble protein 1 6245 CAGCCTTGGAC 3,00 0,91 Hs.65648 RNA binding motif protein 8 6246 ACAGGGTGACC 10,00 2,48 Hs.174050 endothelial differentiation-related factor acid form acid-soluble protein 8 6247 TCCCTGTACAT 5,00 1,40 Hs.89563 nuclear cap binding protein subunit 1,80kD 6248 TTATGGATCTC 5,00 1,40 Hs.5662 guanine nucleotide binding protein (Gprotein), beta 6249 ACTCTGCCAAG 2,00 0,65 Hs.241543 DKFZP586F1524 protein 6250 CAAATGAGGAG 14,00 3,32 Hs.69855 NRAS-related gene 6251 TTCCTCGGGCA 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback						factor 1
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GCTTGCTGAC 1,00 0,32 Hs.1369 decay accelerating factor for complem (CD55, Crom 6243 GATTACCTGTG 1,00 0,32 Hs.119403 hexosaminidase A (alpha polypeptide 6244 TCCGTGGTTGG 7,00 1,81 Hs.79516 brain acid-soluble protein 1 6245 CAGCCTTGGAC 3,00 0,91 Hs.65648 RNA binding motif protein 8 6246 ACAGGGTGACC 10,00 2,48 Hs.174050 endothelial differentiation-related factor 6247 TCCCTGTACAT 5,00 1,40 Hs.89563 nuclear cap binding protein subunit 1,80kD guanine nucleotide binding protein (Gprotein), beta 6249 ACTCTGCCAAG 2,00 0,65 Hs.241543 DKFZP586F1524 protein 6250 CAAATGAGGAG 14,00 3,32 Hs.69855 NRAS-related gene 6251 TTCCTCGGGCA 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback	6241	ACAGCCAAGAG	1,00	0,32	Hs.14142	
6244 TCCGTGGTTGG7,001,81 Hs.79516brain acid-soluble protein 16245 CAGCCTTGGAC3,000,91 Hs.65648RNA binding motif protein 86246 ACAGGGTGACC10,002,48 Hs.174050endothelial differentiation-related factor6247 TCCCTGTACAT5,001,40 Hs.89563nuclear cap binding protein subunit 1,80kD6248 TTATGGATCTC5,001,40 Hs.5662guanine nucleotide binding protein (Gprotein), beta6249 ACTCTGCCAAG2,000,65 Hs.241543DKFZP586F1524 protein6250 CAAATGAGGAG14,003,32 Hs.69855NRAS-related gene6251 TTCCTCGGGCA1,000,33 Hs.83081GTP cyclohydrolase I feedback		·		·		decay accelerating factor for complement (CD55, Crom
6244 TCCGTGGTTGG7,001,81 Hs.79516brain acid-soluble protein 16245 CAGCCTTGGAC3,000,91 Hs.65648RNA binding motif protein 86246 ACAGGGTGACC10,002,48 Hs.174050endothelial differentiation-related factor6247 TCCCTGTACAT5,001,40 Hs.89563nuclear cap binding protein subunit 1,80kD6248 TTATGGATCTC5,001,40 Hs.5662guanine nucleotide binding protein (Gprotein), beta6249 ACTCTGCCAAG2,000,65 Hs.241543DKFZP586F1524 protein6250 CAAATGAGGAG14,003,32 Hs.69855NRAS-related gene6251 TTCCTCGGGCA1,000,33 Hs.83081GTP cyclohydrolase I feedback	6243	GATTACCTGTG	1,00	0,32	Hs.119403	hexosaminidase A (alpha polypeptide)
6246ACAGGGTGACC10,002,48Hs.174050endothelial differentiation-related factor6247TCCCTGTACAT5,001,40Hs.89563nuclear cap binding protein subunit 1,80kD6248TTATGGATCTC5,001,40Hs.5662guanine nucleotide binding protein (Gprotein), beta6249ACTCTGCCAAG2,000,65Hs.241543DKFZP586F1524 protein6250CAAATGAGGAG14,003,32Hs.69855NRAS-related gene6251TTCCTCGGGCA1,000,33Hs.83081GTP cyclohydrolase I feedback	6244	TCCGTGGTTGG	7,00	1,81	Hs.79516	brain acid-soluble protein 1
6246ACAGGGTGACC10,002,48Hs.174050endothelial differentiation-related factor6247TCCCTGTACAT5,001,40Hs.89563nuclear cap binding protein subunit 1,80kD6248TTATGGATCTC5,001,40Hs.5662guanine nucleotide binding protein (Gprotein), beta6249ACTCTGCCAAG2,000,65Hs.241543DKFZP586F1524 protein6250CAAATGAGGAG14,003,32Hs.69855NRAS-related gene6251TTCCTCGGGCA1,000,33Hs.83081GTP cyclohydrolase I feedback	6245	CAGCCTTGGAC	3,00			
80kD     6248   TTATGGATCTC   5,00   1,40   Hs.5662   guanine nucleotide binding protein (G protein), beta   6249   ACTCTGCCAAG   2,00   0,65   Hs.241543   DKFZP586F1524 protein   6250   CAAATGAGGAG   14,00   3,32   Hs.69855   NRAS-related gene   6251   TTCCTCGGGCA   1,00   0,33   Hs.83081   GTP cyclohydrolase I feedback   6260   GTP cyclohydrolase I feedback   626	6246	ACAGGGTGACC	10,00	. 2,48	Hs.174050	endothelial differentiation-related factor 1
6248TTATGGATCTC5,001,40Hs.5662guanine nucleotide binding protein (G protein), beta6249ACTCTGCCAAG2,000,65Hs.241543DKFZP586F1524 protein6250CAAATGAGGAG14,003,32Hs.69855NRAS-related gene6251TTCCTCGGGCA1,000,33Hs.83081GTP cyclohydrolase I feedback	6247	TCCCTGTACAT	5,00	1,40	Hs.89563	nuclear cap binding protein subunit 1,
6249 ACTCTGCCAAG         2,00         0,65 Hs.241543 DKFZP586F1524 protein           6250 CAAATGAGGAG         14,00         3,32 Hs.69855 NRAS-related gene           6251 TTCCTCGGGCA         1,00         0,33 Hs.83081 GTP cyclohydrolase I feedback	6248	TTATGGATCTC	5,00	1,40	Hs.5662	guanine nucleotide binding protein (G
6250 CAAATGAGGAG 14,00 3,32 Hs.69855 NRAS-related gene 6251 TTCCTCGGGCA 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback	6249	ACTCTGCCAAG	2.00	0.65	Hs.241543	
6251 TTCCTCGGGCA 1,00 0,33 Hs.83081 GTP cyclohydrolase I feedback						
1						
						regulatory protein
6252 TTAAATGCAAA 1,00 0,33 Hs.7935 KIAA0952 protein	6252	TTAAATGCAAA	1,00	0,33	Hs.7935	<del>                                     </del>

S253   TCTGGACCGGC   1,00   0,33   Hs. 78979   Golgi apparatus protein 1					<del></del>	<del></del>
6255         ACCTTATCAAC         1,00         0,33 Hs.75659         MpV17 transgene, murine homolog, glomerulosclerosis           6256         CCTGGGGGTGC         1,00         0,33 Hs.75196         ankyrin repeat-containing protein           6257         ACAGCGTCTGC         1,00         0,33 Hs.63128         KIAA1292 protein           6258         GCACAGAGCCG         1,00         0,33 Hs.25126         RAB9, member RAS oncogene family           6260         GTTGTGCTTT         1,00         0,33 Hs.279822         PC326 protein           6261         GATTTCTACT         1,00         0,33 Hs.17969         KIAA0663 gene product           6261         GATTTCTACT         1,00         0,33 Hs.11355         ESTs           6263         TACTTGGTCTT         1,00         0,33 Hs.11355         ESTs           6263         TACTTGGTCTT         1,00         0,33 Hs.11355         ESTs           6264         TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT			1,00			Golgi apparatus protein 1
glomerulosclerosis   glomeru						
6256   CCTGGGGGTGC         1,00         0,33   Hs. 75196         Inklyrin repeat-containing protein           6257   ACAGCGTCTGC         1,00         0,33   Hs. 63128         KIAA1292 protein           6258   GCACAGAGCCG         1,00         0,33   Hs. 28726         RAB9, member RAS oncogene family           6259   GGATGGAGAA         1,00         0,33   Hs. 279882   PC326 protein           6261   GATTTCTTCT         1,00         0,33   Hs. 13014         ADP-ribosylation factor GTPase activating protein 1           6262   GGCTCAGGGCT         1,00         0,33   Hs. 11355   ESTs         ESCAS   TACTTGGTCTT           6263   TACTTGGTCTT         1,00         0,33   Hs. 112058   CD27-binding (Siva) protein           6264   TTCTATTTGT         1,00         0,33   Hs. 179735   ras homolog gene family, member C           6265   CATGTGATGC         16,00         3,77   Hs. 249495   heterogeneous nuclear ribonucleoprotein A1           6266   CTGTTGATTGC         16,00         1,66   Hs. 3100   lysyl-RNA synthetase           6267   AAACCCGAAGA         2,00         0,66   Hs. 3100   lysyl-RNA synthetase           6268   GAAATTTAAAG         6,00         1,66   Hs. 274472   lymin hybridity group (nonhistone chromosomal) protein           6270   AAGGGAGGTGT         4,00         1,20   Hs. 77886   lamin A/C           6271   GTCACCTCCTG         1,00         0,34   Hs. 182248   sequestoso	6255	ACCTTATCAAC	1,00	0,33	Hs.75659	
6257   ACAGCGTCTGC         1,00         0,33   Hs. 63128   KIAA1292 protein           6258   GCACAGAGCG         1,00         0,33   Hs. 5105   hypothetical protein FLJ10569           6259   GGATGGAGAA   1,00   0,33   Hs. 28726   RAB9, member RAS oncogene family         6260   TTTGTTGCTTT   1,00   0,33   Hs. 279882   PC326 protein           6261   GATTTTCTACT   1,00   0,33   Hs. 17969   KIAA0663 gene product         6262   GGCTCAGGGCT   1,00   0,33   Hs. 1301   ADP-ribosylation factor GTPase activating protein 1           6263   TACTTGGTCTT   1,00   0,33   Hs. 11305   ESTs   6264   TTCATTTTGT   1,00   0,33   Hs. 112058   CD27-binding (Siva) protein           6265   AATAAAGGCTA   7,00   1,86   Hs. 179735   ras homolog gene family, member C           6266   CTGTTGATTGC   16,00   3,77   Hs. 249495   heterogeneous nuclear ribonucleoprotein   A1           6267   AAACCCGAAGA   2,00   0,66   Hs. 3100   lysyl-tRNA synthetase   6268   GAAATTTAAAG   6,00   1,66   Hs. 274472   high-mobility group (nonhistone chromosomal) protein           6269   CGCAAGCTGGT   4,00   1,20   Hs. 77886   lamin A/C   6270   AAGGGAGGGTC   5,00   1,44   Hs. 182248   sequestosome 1   6271   GTCTACTCCTC   1,00   0,34   Hs. 7345   MAD1 (mitotic arrest deficient, yeast, homolog)-like   homo sapiens mRNA; cDNA   DKF2,761E1423 (from clone DK   6274   CTTATTCCTTA   1,00   0,34   Hs. 62771   Homo sapiens mRNA; cDNA   DKF2,761E1423 (from clone DK   6276   CACTGCAAGGC   1,00   0,34   Hs. 283109   hypothetical protein DKF2,762L1710   6276   CACTGCAAGCC   1,00   0,34   Hs. 283109   hypothetical protein DKF2,762L1710   6283   GGCAACACCTG   1,00   0,34   Hs. 18894   adaptor-related protein complex 1, mu 2 subunit   6280   AGTGACCAACCTG   1,00   0,34   Hs. 134342   TASP for testis						
6258 GCACAGAGCCG         1,00         0,33 Hs.5105         hypothetical protein FLJ10569           6259 GGGATGGAGAA         1,00         0,33 Hs.28726         RAB9, member RAS oncogene family           6260 TTTGTTGCTTT         1,00         0,33 Hs.27982 PC326 protein           6261 GATTTCTACT         1,00         0,33 Hs.17969         KIAA0683 gene product           6262 GGCTCAGGGCT         1,00         0,33 Hs.13014         ADP-ribosylation factor GTPase activating protein           6263 TACTTGTCTT         1,00         0,33 Hs.113055         ESTs           6264 ATACTATITTGT         1,00         0,33 Hs.112058 (CD27-binding (Siva) protein           6265 ATAAAGGCTA         7,00         1,86 Hs.179735 ras homolog gene family, member C           6266 CTGTTGATTGC         16,00         3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1           6267 AAACCCGAAGA         2,00         0,66 Hs.3100         lysyl-IRNA synthetase           6268 GAAATTTAAAG         6,00         1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein           6270 AAAGGAGAGGCTG         4,00         1,20 Hs.77886         lamin A/C           6271 GTCACCCGGT         4,00         1,24 Hs.75199         protein phosphatase 2, regulatory subunit B (B56), b           6272 CGTCCCGGAGC         1,00         0,34 Hs.62771         Homo sapiens mRNA; cD	6256	CCTGGGGGTGC	1,00	0,33	Hs.75196	
8259 GGGATGGAGAA   1,00   0,33   Hs.28726   RAB9, member RAS oncogene family   9260 TTTGTTGCTTT   1,00   0,33   Hs.27882   PC326 protein   9261 GATTTTCTACT   1,00   0,33   Hs.17989   KIAA0663 gene product   9261 GATTTTCTACT   1,00   0,33   Hs.17989   KIAA0663 gene product   9263 TACTTGGTCTT   1,00   0,33   Hs.13014   ADP-ribosylation factor GTPase activating protein   9263 TACTTGGTCTT   1,00   0,33   Hs.13051   ESTS   9264 TTCTATTTTGT   1,00   0,33   Hs.112058   CD27-binding (Siva) protein   9265 AATAAAGGCTA   7,00   1,86   Hs.179735  ras homolog gene family, member C   9266 CTGTTGATTGC   16,00   3,77   Hs.249495   heterogeneous nuclear ribonucleoprotein   A1   9266 CGAAGACACACACACACACACACACACACACACACACAC	6257	ACAGCGTCTGC		0,33	Hs.63128	KIAA1292 protein
6260   TTTGTTGCTTT	6258	GCACAGAGCCG	1,00	0,33	Hs.5105	hypothetical protein FLJ10569
6281 GATTTCTACT         1,00         0,33 Hs.17969         KIAA0663 gene product           6282 GGCTCAGGGCT         1,00         0,33 Hs.13014         ADP-ribosylation factor GTPase activating protein 1           6283 TACTTGGTCTT         1,00         0,33 Hs.11355         ESTs           6264 TTCTATTTTGT         1,00         0,33 Hs.112058 CD27-binding (Siva) protein           6265 AATAAAGGCTA         7,00         1,86 Hs.179735 ras homolog gene family, member C           6266 CTGTTGATTGC         16,00         3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1           6267 AAACCCGAAGA         2,00         0,66 Hs.3100         lysyl-tRNA synthetase           6268 GAAATTTAAAG         6,00         1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein           6269 GCCAAGCTGGT         4,00         1,20 Hs.77886 lamin A/C           6270 AAGGGAGGGTC         5,00         1,44 Hs.182248 sequestosome 1           6271 GTCTACTCCTC         1,00         0,34 Hs.7345         MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273 GGACCAGGCTG         1,00         0,34 Hs.62771         Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274 CTTATTCCTTA         1,00         0,34 Hs.20323         BUB3 (budding unlinhibited by benzimidazoles 3, yeast           62776 CACTGCAAGGC         1,00         0,34 Hs.3040	6259	GGGATGGAGAA	1,00	0,33	Hs.28726	RAB9, member RAS oncogene family
6262         GGCTCAGGGCT         1,00         0,33 Hs.13014         ADP-ribosylation factor GTPase activating protein 1           6263         TACTTGGTCTT         1,00         0,33 Hs.11355         ESTs           6264         TTCTATTTTGT         1,00         0,33 Hs.112058         CD27-binding (Siva) protein           6265         CATAAAGGCTA         7,00         1,86 Hs.179735 ras homolog gene family, member C           6266         CTGTTGATTGC         16,00         3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1           6267         AAACCCGAAGA         2,00         0,66 Hs.3100         lysyl-tRNA synthetase           6268         GAAATTTAAAG         6,00         1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein           6269         CGCAAGCTGGT         4,00         1,20 Hs.77886         lamin A/C           6270         AAGGGAGGGTC         5,00         1,44 Hs.182248 sequestosome 1           6271         GTCTACTCCTC         1,00         0,34 Hs.7345         MAD1 (mitotic arrest deficient, yeast, homolog)-like           6272         CGTCCCGGAGC         1,00         0,34 Hs.62771         Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274         CTTATTCCTTA         1,00         0,34 Hs.32060         ESTs           6275         CACTGCAAGGC<	6260	TTTGTTGCTTT	1,00	0,33	Hs.279882	PC326 protein
activating protein 1	6261	GATTTTCTACT	1,00	0,33	Hs.17969	KIAA0663 gene product
6263 TACTTGGTCTT         1,00         0,33 Hs.11355         ESTs           6264 TTCTATTTTGT         1,00         0,33 Hs.112058 CD27-binding (Siva) protein           6265 AATAAAGGCTA         7,00         1,86 Hs.179735 ras homolog gene family, member C           6266 CTGTTGATTGC         16,00         3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1           6267 AAACCCGAAGA         2,00         0,66 Hs.3100         lysyl-tRNA synthetase           6268 GAAATTTAAAG         6,00         1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein           6269 CGCAAGCTGGT         4,00         1,20 Hs.77886 lamin A/C           6270 AAGGGAGGGTC         5,00         1,44 Hs.182248 sequestosome 1           6271 GTCACTCCTC         1,00         0,34 Hs.7345 protein phosphatase 2, regulatory subunit B (B56), b           6272 CGTCCCGGAGC         1,00         0,34 Hs.62771 homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6273 GGACCAGGCTG         1,00         0,34 Hs.40323 bUB3 (budding uninhibited by benzimidazoles 3, yeast           6275 CACTGCAAGGC         1,00         0,34 Hs.283109 hypothetical protein DKFZp762L1710           6275 CACTGCAAGGC         1,00         0,34 Hs.283109 hypothetical protein complex 1, mu 2 subunit           6276 GCAACACATCT         1,00         0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit           627	6262	GGCTCAGGGCT	1,00	0,33	Hs.13014	ADP-ribosylation factor GTPase
6264 TTCTATTTGT         1,00         0,33 Hs.112058 CD27-binding (Siva) protein           6265 AATAAAAGGCTA         7,00         1,86 Hs.179735 ras homolog gene family, member C           6266 CTGTTGATTGC         16,00         3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1           6267 AAACCCGAAGA         2,00         0,66 Hs.3100 lysyl-tRNA synthetase           6268 GAAATTTAAAG         6,00         1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein           6269 CGCAAGCTGGT         4,00         1,20 Hs.77886 lamin A/C           6270 AAGGGAGGGTC         5,00         1,44 Hs.182248 sequestosome 1           6271 GTCTACTCCTC         1,00         0,34 Hs.7345 mclosup protein phosphatase 2, regulatory subunit 8 (B56), b           6272 CGTCCCGGAGC         1,00         0,34 Hs.62771 mclosup protein phosphatase 2, regulatory subunit 8 (B56), b           6273 GGACCAGGCTG         1,00         0,34 Hs.62771 mclosup protein phosphatase 2, regulatory subunit 8 (B56), b           6274 CTTATTCCTTA         1,00         0,34 Hs.62771 mclosup protein phosphatase 2, regulatory subunit 8 (B56), b           6275 CACTGCAGGCC         1,00         0,34 Hs.40323 mclosup protein phosphatase 2, regulatory subunit 8 (B56), b           6275 CACTGCAGGCC         1,00         0,34 Hs.32060 mclosup protein phosphatase 2, regulatory subunit 8 (B56), b           6275 CACTGCAAGGC         1,00         0,34 Hs.32060						activating protein 1
6265 AATAAAGGCTA 7,00 1,86 Hs.179735 ras homolog gene family, member C 6266 CTGTTGATTGC 16,00 3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1 6267 AAACCCGAAGA 2,00 0,66 Hs.3100 lysyl-tRNA synthetase 6268 GAAATTTAAAG 6,00 1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein 6269 CGCAAGCTGGT 4,00 1,20 Hs.77886 lamin A/C 6270 AAGGGAGGGTC 5,00 1,44 Hs.182248 sequestosome 1 6271 GTCTACTCCTC 1,00 0,34 Hs.75199 protein phosphatase 2, regulatory subunit B (B56), b 6272 CGTCCCGGAGC 1,00 0,34 Hs.62771 homosapiens mRNA; cDNA 6274 CTTATTCCTTA 1,00 0,34 Hs.40323 BUB3 (budding uninhibited by benzimidazoles 3, yeast 6275 CACTGCAAGGC 1,00 0,34 Hs.32060 ESTs 6276 GCAACACATCT 1,00 0,34 Hs.19165 ESTs 6276 GCAACACATCAT 1,00 0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit 6279 GTCCCGAAG 1,00 0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit 6280 AGTGACCGAAG 1,00 0,34 Hs.19165 ESTs 6281 TAATTTTCTA 1,00 0,34 Hs.193432 TASP for testis-specific adriamycin sensitivity prot 6281 TAATTTTCTA 1,00 0,34 Hs.197637 Homo sapiens clone 25107 mRNA 6282 CAGTTCTCGG 4,00 1,20 Hs.19349 ESTs 6283 GGCTGGTCTG 4,00 1,20 Hs.7891 Hs.9C035 protein 6284 AGCCCTCCTG 6,00 1,69 Hs.74111 RNA-binding protein (autoantigenic) 6285 CAGGCTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAACCTA 2,00 0,67 Hs.8799 DEAD/H (Asp-Glu-Ala-Asp/His) box	6263	TACTTGGTCTT	1,00			
6265 AATAAAGGCTA 7,00 1,86 Hs.179735 ras homolog gene family, member C 6266 CTGTTGATTGC 16,00 3,77 Hs.249495 heterogeneous nuclear ribonucleoprotein A1 6267 AAACCCGAAGA 2,00 0,66 Hs.3100 lysyl-tRNA synthetase 6268 GAAATTTAAAG 6,00 1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein 6269 CGCAAGCTGGT 4,00 1,20 Hs.77886 lamin A/C 6270 AAGGGAGGGTC 5,00 1,44 Hs.182248 sequestosome 1 6271 GTCTACTCCTC 1,00 0,34 Hs.75199 protein phosphatase 2, regulatory subunit B (B56), b 6272 CGTCCCGGAGC 1,00 0,34 Hs.62771 homosapiens mRNA; cDNA 6274 CTTATTCCTTA 1,00 0,34 Hs.40323 BUB3 (budding uninhibited by benzimidazoles 3, yeast 6275 CACTGCAAGGC 1,00 0,34 Hs.32060 ESTs 6276 GCAACACATCT 1,00 0,34 Hs.19165 ESTs 6276 GCAACACATCAT 1,00 0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit 6279 GTCCCGAAG 1,00 0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit 6280 AGTGACCGAAG 1,00 0,34 Hs.19165 ESTs 6281 TAATTTTCTA 1,00 0,34 Hs.193432 TASP for testis-specific adriamycin sensitivity prot 6281 TAATTTTCTA 1,00 0,34 Hs.197637 Homo sapiens clone 25107 mRNA 6282 CAGTTCTCGG 4,00 1,20 Hs.19349 ESTs 6283 GGCTGGTCTG 4,00 1,20 Hs.7891 Hs.9C035 protein 6284 AGCCCTCCTG 6,00 1,69 Hs.74111 RNA-binding protein (autoantigenic) 6285 CAGGCTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAACCTA 2,00 0,67 Hs.8799 DEAD/H (Asp-Glu-Ala-Asp/His) box	6264	TTCTATTTTGT	1,00	0,33	Hs.112058	CD27-binding (Siva) protein
AAACCCGAAGA   2,00   0,66   Hs.3100    ysyl-tRNA synthetase   6268   GAAATTTAAAG   6,00   1,66   Hs.274472   high-mobility group (nonhistone chromosomal) protein   6269   CGCAAGCTGGT   4,00   1,20   Hs.77886   lamin A/C   6270   AAGGGAGGGTC   5,00   1,44   Hs.182248   sequestosome 1   6271   GTCTACTCCTC   1,00   0,34   Hs.75199   protein phosphatase 2, regulatory subunit B (B56), b   MAD1 (mitotic arrest deficient, yeast, homolog)-like   6272   CGTCCCGGAGC   1,00   0,34   Hs.62771   Horno sapiens mRNA; cDNA   DKFZp761E1423 (from clone DK   6274   CTTATTCCTTA   1,00   0,34   Hs.40323   BUB3 (budding uninhibited by benzimidazoles 3, yeast   6275   CACTGCAAGGC   1,00   0,34   Hs.283109   hypothetical protein DKFZp762L1710   6277   GCTTCCGGCCC   1,00   0,34   Hs.19165   ESTs   6278   AAGATCATTGA   1,00   0,34   Hs.18894   adaptor-related protein complex 1, mu 2   subunit   6280   AGTGACCGAAG   1,00   0,34   Hs.159608   aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen   6280   AGTGACCGAAG   1,00   0,34   Hs.134342   TASP for testis-specific adriamycin   sensitivity prot   6281   TAATTTTCTA   1,00   0,34   Hs.134342   TASP for testis-specific adriamycin   sensitivity prot   6282   CAGTTCTCTGT   7,00   1,91   Hs.279921   HSPC035 protein   6283   GCCTGCTCG   6,00   1,69   Hs.74111   RNA-binding protein (autoantigenic)   6285   CAGGCTTTTTG   2,00   0,67   Hs.83484   SRY (sex determining region Y)-box 4   6286   GAAAAGCCTTC   2,00   0,67   Hs.6179   DEAD/H (Asp-Glu-Ala-Asp/His) box	6265	AATAAAGGCTA	7,00			
6267         AAACCCGAAGA         2,00         0,66         Hs.3100         lysyl-tRNA synthetase           6268         GAAATTTAAAG         6,00         1,66         Hs.274472         high-mobility group (nonhistone chromosomal) protein           6269         CGCAAGCTGGT         4,00         1,20         Hs.77886         lamin A/C           6270         AAGGGAGGGTC         5,00         1,44         Hs.182248 sequestosome 1           6271         GTCTACTCCTC         1,00         0,34         Hs.75199         protein phosphatase 2, regulatory subunit B (B56), b           6272         CGTCCCGGAGC         1,00         0,34         Hs.7345         MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273         GGACCAGGCTG         1,00         0,34         Hs.62771         Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274         CTTATTCCTTA         1,00         0,34         Hs.32060         ESTs           6276         CAACGAAGCT         1,00         0,34         Hs.283109 hypothetical protein DKFZp762L1710           6277         GCTTCCGGCCC         1,00         0,34         Hs.19165         ESTs           6278         AAGATCATTGA         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit	6266	CTGTTGATTGC	16,00			
6268 GAAATTTAAAG 6,00 1,66 Hs.274472 high-mobility group (nonhistone chromosomal) protein lamin A/C 6270 AAGGGAGGTC 5,00 1,44 Hs.182248 sequestosome 1 6271 GTCTACTCCTC 1,00 0,34 Hs.75199 protein phosphatase 2, regulatory subunit B (B56), b 6272 CGTCCCGGAGC 1,00 0,34 Hs.7345 MAD1 (mitotic arrest deficient, yeast, homolog)-like 6273 GGACCAGGCTG 1,00 0,34 Hs.62771 Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK 6274 CTTATTCCTTA 1,00 0,34 Hs.40323 BUB3 (budding uninhibited by benzimidazoles 3, yeast 6275 CACTGCAAGGC 1,00 0,34 Hs.283109 hypothetical protein DKFZp762L1710 6277 GCTTCCGGCCC 1,00 0,34 Hs.19165 ESTs 6278 AAGATCATTGA 1,00 0,34 Hs.19894 adaptor-related protein complex 1, mu 2 subunit 6280 AGTGACCAAGC 1,00 0,34 Hs.19608 aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen 6280 AGTGACCGAAG 1,00 0,34 Hs.19608 aldehyde dehydrogen 6280 AGTGACCAAGCCTCC 1,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTC 2,00 0,67 Hs.8619 DEAD/H (Asp-Glu-Ala-Asp/His) box						A1
Chromosomal	6267	AAACCCGAAGA	2,00	0,66	Hs.3100	lysyl-tRNA synthetase
6269 CGCAAGCTGGT         4,00         1,20 Hs.77886         lamin A/C           6270 AAGGGAGGGTC         5,00         1,44 Hs.182248 sequestosome 1           6271 GTCTACTCCTC         1,00         0,34 Hs.75199 protein phosphatase 2, regulatory subunit B (B56), b           6272 CGTCCCGGAGC         1,00         0,34 Hs.7345 MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273 GGACCAGGCTG         1,00         0,34 Hs.62771 Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274 CTTATTCCTTA         1,00         0,34 Hs.40323 BUB3 (budding uninhibited by benzimidazoles 3, yeast           6275 CACTGCAAGGC         1,00         0,34 Hs.32060 ESTs           6276 GCAACACATCT         1,00         0,34 Hs.18309 hypothetical protein DKFZp762L1710           6277 GCTTCCGGCCC         1,00         0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit           6278 AAGATCATTGA         1,00         0,34 Hs.1894 aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6280 AGTGACCGAAG         1,00         0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot           6281 TAATTTTCTA         1,00         0,34 Hs.107637 Hs.107637 Homo sapiens clone 25107 mRNA sequence           6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GCCGGCCTGGG 6,00         1,69 Hs.74111 RNA-binding protein (autoantigenic)           6285 C	6268	GAAATTTAAAG	6,00	1,66	Hs.274472	high-mobility group (nonhistone
6270 AAGGGAGGGTC         5,00         1,44 Hs. 182248 sequestosome 1           6271 GTCTACTCCTC         1,00         0,34 Hs. 75199 protein phosphatase 2, regulatory subunit B (B56), b           6272 CGTCCCGGAGC         1,00         0,34 Hs. 7345 MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273 GGACCAGGCTG         1,00         0,34 Hs. 62771 Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274 CTTATTCCTTA         1,00         0,34 Hs. 40323 BUB3 (budding uninhibited by benzimidazoles 3, yeast           6275 CACTGCAAGGC         1,00         0,34 Hs. 283109 hypothetical protein DKFZp762L1710           6276 GCAACACATCT         1,00         0,34 Hs. 19165 ESTs           6278 AAGATCATTGA         1,00         0,34 Hs. 18894 adaptor-related protein complex 1, mu 2 subunit           6279 GTCACAACCTG         1,00         0,34 Hs. 159608 aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6280 AGTGACCGAAG         1,00         0,34 Hs. 134342 TASP for testis-specific adriamycin sensitivity prot           6281 TAATTTTCTA         1,00         0,34 Hs. 107637 Homo sapiens clone 25107 mRNA sequence           6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GCTGGTCTGG         4,00         1,20 Hs. 17894 ESTs           6284 AGCCCTCCCTG         6,00         1,69 Hs. 74111 RNA-binding protein (autoantigenic)           6285 C					·	chromosomal) protein
6271         GTCTACTCCTC         1,00         0,34         Hs.75199         protein phosphatase 2, regulatory subunit B (B56), b           6272         CGTCCCGGAGC         1,00         0,34         Hs.7345         MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273         GGACCAGGCTG         1,00         0,34         Hs.62771         Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274         CTTATTCCTTA         1,00         0,34         Hs.40323         BUB3 (budding uninhibited by benzimidazoles 3, yeast           6275         CACTGCAAGGC         1,00         0,34         Hs.283109 hypothetical protein DKFZp762L1710           6276         GCAACACATCT         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6279         GTCACAACCTG         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6280         AGTGACCGAAG         1,00         0,34         Hs.159608         aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6281         TAATTTTCTA         1,00         0,34         Hs.134342         TASP for testis-specific adriamycin sensitivity prot           6281         TAATTTTCTA         1,00         0,34         Hs.279921         HSPC035 protein           6282	6269	CGCAAGCTGGT	4,00	1,20	Hs.77886	lamin A/C
6271         GTCTACTCCTC         1,00         0,34         Hs.75199         protein phosphatase 2, regulatory subunit B (B56), b           6272         CGTCCCGGAGC         1,00         0,34         Hs.7345         MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273         GGACCAGGCTG         1,00         0,34         Hs.62771         Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274         CTTATTCCTTA         1,00         0,34         Hs.40323         BUB3 (budding uninhibited by benzimidazoles 3, yeast           6275         CACTGCAAGGC         1,00         0,34         Hs.283109 hypothetical protein DKFZp762L1710           6276         GCAACACATCT         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6279         GTCACAACCTG         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6280         AGTGACCGAAG         1,00         0,34         Hs.159608         aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6281         TAATTTTCTA         1,00         0,34         Hs.134342         TASP for testis-specific adriamycin sensitivity prot           6281         TAATTTTCTA         1,00         0,34         Hs.279921         HSPC035 protein           6282	6270	AAGGGAGGGTC	5,00	1,44	Hs.182248	sequestosome 1
Subunit B (B56), b	6271	GTCTACTCCTC	1,00			
6272         CGTCCCGGAGC         1,00         0,34 Hs.7345         MAD1 (mitotic arrest deficient, yeast, homolog)-like           6273         GGACCAGGCTG         1,00         0,34 Hs.62771         Homo sapiens mRNA; cDNA DKFZp761E1423 (from clone DK           6274         CTTATTCCTTA         1,00         0,34 Hs.40323         BUB3 (budding uninhibited by benzimidazoles 3, yeast           6275         CACTGCAAGGC         1,00         0,34 Hs.32060         ESTs           6276         GCAACACATCT         1,00         0,34 Hs.283109 hypothetical protein DKFZp762L1710           6277         GCTTCCGGCCC         1,00         0,34 Hs.19165         ESTs           6278         AAGATCATTGA         1,00         0,34 Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6280         AGTGACCGAAG         1,00         0,34 Hs.159608         aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6280         AGTGACCGAAG         1,00         0,34 Hs.134342         TASP for testis-specific adriamycin sensitivity prot           6281         TAATTTTCTA         1,00         0,34 Hs.107637         Homo sapiens clone 25107 mRNA sequence           6282         CAGTCCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283         GGCTGGTCTGG         4,00         1,20 Hs.17894         <			}			
homolog)-like   homology-like   homology-lik	6272	CGTCCCGGAGC	1,00	0,34	Hs.7345	
DKFZp761E1423 (from clone DK						
6274         CTTATTCCTTA         1,00         0,34         Hs.40323         BUB3 (budding uninhibited by benzimidazoles 3, yeast           6275         CACTGCAAGGC         1,00         0,34         Hs.32060         ESTs           6276         GCAACACATCT         1,00         0,34         Hs.283109 hypothetical protein DKFZp762L1710           6277         GCTTCCGGCCC         1,00         0,34         Hs.19165         ESTs           6278         AAGATCATTGA         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6279         GTCACAACCTG         1,00         0,34         Hs.159608         aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6280         AGTGACCGAAG         1,00         0,34         Hs.134342         TASP for testis-specific adriamycin sensitivity prot           6281         TAATTTTTCTA         1,00         0,34         Hs.107637         Homo sapiens clone 25107 mRNA sequence           6282         CAGTTCTCTGT         7,00         1,91         Hs.279921         HSPC035 protein           6283         GGCTGGTCTGG         4,00         1,20         Hs.17894         ESTs           6284         AGCCCTCCCTG         6,00         1,69         Hs.74111         RNA-binding protein (autoantigen	6273	GGACCAGGCTG	1,00	0,34	Hs.62771	Homo sapiens mRNA; cDNA
benzimidazoles 3, yeast						DKFZp761E1423 (from clone DK
6275 CACTGCAAGGC         1,00         0,34 Hs.32060         ESTs           6276 GCAACACATCT         1,00         0,34 Hs.283109 hypothetical protein DKFZp762L1710           6277 GCTTCCGGCCC         1,00         0,34 Hs.19165         ESTs           6278 AAGATCATTGA         1,00         0,34 Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6279 GTCACAACCTG         1,00         0,34 Hs.159608         aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6280 AGTGACCGAAG         1,00         0,34 Hs.134342         TASP for testis-specific adriamycin sensitivity prot           6281 TAATTTTTCTA         1,00         0,34 Hs.107637         Homo sapiens clone 25107 mRNA sequence           6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GGCTGGTCTGG         4,00         1,20 Hs.17894         ESTs           6284 AGCCCTCCCTG         6,00         1,69 Hs.74111         RNA-binding protein (autoantigenic)           6285 CAGGCTTTTTG         2,00         0,67 Hs.83484         SRY (sex determining region Y)-box 4           6286 GAAAAGCCTTC         2,00         0,67 Hs.86179         DEAD/H (Asp-Glu-Ala-Asp/His) box	6274	CTTATTCCTTA	1,00	0,34	Hs.40323	
6276         GCAACACATCT         1,00         0,34         Hs.283109         hypothetical protein DKFZp762L1710           6277         GCTTCCGGCCC         1,00         0,34         Hs.19165         ESTs           6278         AAGATCATTGA         1,00         0,34         Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6279         GTCACAACCTG         1,00         0,34         Hs.159608         aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen           6280         AGTGACCGAAG         1,00         0,34         Hs.134342         TASP for testis-specific adriamycin sensitivity prot           6281         TAATTTTCTA         1,00         0,34         Hs.107637         Homo sapiens clone 25107 mRNA sequence           6282         CAGTTCTCTGT         7,00         1,91         Hs.279921         HSPC035 protein           6283         GGCTGGTCTGG         4,00         1,20         Hs.17894         ESTs           6284         AGCCCTCCCTG         6,00         1,69         Hs.74111         RNA-binding protein (autoantigenic)           6285         CAGGCTTTTTG         2,00         0,67         Hs.83484         SRY (sex determining region Y)-box 4           6286         GAAAAGCCTTC         2,00         0,67         Hs.6179         DE						
6277 GCTTCCGGCCC         1,00         0,34 Hs.19165         ESTs           6278 AAGATCATTGA         1,00         0,34 Hs.18894         adaptor-related protein complex 1, mu 2 subunit           6279 GTCACAACCTG         1,00         0,34 Hs.159608 aldehyde dehydrogenses 10 (fatty aldehyde dehydrogen           6280 AGTGACCGAAG         1,00         0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot           6281 TAATTTTTCTA         1,00         0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence           6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GGCTGGTCTGG         4,00         1,20 Hs.17894 ESTs           6284 AGCCCTCCCTG         6,00         1,69 Hs.74111 RNA-binding protein (autoantigenic)           6285 CAGGCTTTTTG         2,00         0,67 Hs.83484 SRY (sex determining region Y)-box 4           6286 GAAAAGCCTTC         2,00         0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag           6287 GTTGAAACTCA         2,00         0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box						
6278 AAGATCATTGA  1,00  0,34 Hs.18894 adaptor-related protein complex 1, mu 2 subunit  6279 GTCACAACCTG  1,00  0,34 Hs.159608 aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen  6280 AGTGACCGAAG  1,00  0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot  6281 TAATTTTTCTA  1,00  0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence  6282 CAGTTCTCTGT  7,00  1,91 Hs.279921 HSPC035 protein  6283 GGCTGGTCTGG  4,00  1,20 Hs.17894 ESTs  6284 AGCCCTCCCTG  6,00  1,69 Hs.74111 RNA-binding protein (autoantigenic)  6285 CAGGCTTTTTG  2,00  0,67 Hs.83484 SRY (sex determining region Y)-box 4  6286 GAAAAGCCTTC  2,00  0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag  6287 GTTGAAACTCA  2,00  0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box				0,34	Hs.283109	hypothetical protein DKFZp762L1710
subunit 6279 GTCACAACCTG 1,00 0,34 Hs.159608 aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen 6280 AGTGACCGAAG 1,00 0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot 6281 TAATTTTCTA 1,00 0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence 6282 CAGTTCTCTGT 7,00 1,91 Hs.279921 HSPC035 protein 6283 GGCTGGTCTGG 4,00 1,20 Hs.17894 ESTs 6284 AGCCCTCCCTG 6,00 1,69 Hs.74111 RNA-binding protein (autoantigenic) 6285 CAGGCTTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box			1,00	0,34	Hs.19165	ESTs
GTCACAACCTG 1,00 0,34 Hs.159608 aldehyde dehydrogenase 10 (fatty aldehyde dehydrogen 6280 AGTGACCGAAG 1,00 0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot 6281 TAATTTTCTA 1,00 0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence 6282 CAGTTCTCTGT 7,00 1,91 Hs.279921 HSPC035 protein 6283 GGCTGGTCTGG 4,00 1,20 Hs.17894 ESTs 6284 AGCCCTCCCTG 6,00 1,69 Hs.74111 RNA-binding protein (autoantigenic) 6285 CAGGCTTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box	6278	AAGATCATTGA	1,00	0,34	Hs.18894	adaptor-related protein complex 1, mu 2
aldehyde dehydrogen  6280 AGTGACCGAAG  1,00  0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot  6281 TAATTTTCTA  1,00  0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence  6282 CAGTTCTCTGT  7,00  1,91 Hs.279921 HSPC035 protein  6283 GGCTGGTCTGG  4,00  1,20 Hs.17894 ESTs  6284 AGCCCTCCCTG  6,00  1,69 Hs.74111 RNA-binding protein (autoantigenic)  6285 CAGGCTTTTTG  2,00  0,67 Hs.83484 SRY (sex determining region Y)-box 4  6286 GAAAAGCCTTC  2,00  0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag  6287 GTTGAAACTCA  2,00  0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box	· ·					
aldehyde dehydrogen  6280 AGTGACCGAAG  1,00  0,34 Hs.134342 TASP for testis-specific adriamycin sensitivity prot  6281 TAATTTTCTA  1,00  0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence  6282 CAGTTCTCTGT  7,00  1,91 Hs.279921 HSPC035 protein  6283 GGCTGGTCTGG  4,00  1,20 Hs.17894 ESTs  6284 AGCCCTCCCTG  6,00  1,69 Hs.74111 RNA-binding protein (autoantigenic)  6285 CAGGCTTTTTG  2,00  0,67 Hs.83484 SRY (sex determining region Y)-box 4  6286 GAAAAGCCTTC  2,00  0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag  6287 GTTGAAACTCA  2,00  0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box	6279	GTCACAACCTG	1,00	0,34	Hs.159608	aldehyde dehydrogenase 10 (fatty
Sensitivity prot						aldehyde dehydrogen
6281 TAATTTTCTA         1,00         0,34 Hs.107637 Homo sapiens clone 25107 mRNA sequence           6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GGCTGGTCTGG         4,00         1,20 Hs.17894 ESTs           6284 AGCCCTCCCTG         6,00         1,69 Hs.74111 RNA-binding protein (autoantigenic)           6285 CAGGCTTTTTG         2,00         0,67 Hs.83484 SRY (sex determining region Y)-box 4           6286 GAAAAGCCTTC         2,00         0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag           6287 GTTGAAACTCA         2,00         0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box	6280	AGTGACCGAAG	1,00	0,34	Hs.134342	
6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GGCTGGTCTGG         4,00         1,20 Hs.17894 ESTs           6284 AGCCCTCCCTG         6,00         1,69 Hs.74111 RNA-binding protein (autoantigenic)           6285 CAGGCTTTTTG         2,00         0,67 Hs.83484 SRY (sex determining region Y)-box 4           6286 GAAAAGCCTTC         2,00         0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag           6287 GTTGAAACTCA         2,00         0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box		<u> </u>				
6282 CAGTTCTCTGT         7,00         1,91 Hs.279921 HSPC035 protein           6283 GGCTGGTCTGG         4,00         1,20 Hs.17894 ESTs           6284 AGCCCTCCCTG         6,00         1,69 Hs.74111 RNA-binding protein (autoantigenic)           6285 CAGGCTTTTTG         2,00         0,67 Hs.83484 SRY (sex determining region Y)-box 4           6286 GAAAAGCCTTC         2,00         0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag           6287 GTTGAAACTCA         2,00         0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box	6281	TAATTTTTCTA	1,00	0,34	Hs.107637	Homo sapiens clone 25107 mRNA
6283 GGCTGGTCTGG 4,00 1,20 Hs.17894 ESTs 6284 AGCCCTCCCTG 6,00 1,69 Hs.74111 RNA-binding protein (autoantigenic) 6285 CAGGCTTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box						
6284 AGCCCTCCTG 6,00 1,69 Hs.74111 RNA-binding protein (autoantigenic) 6285 CAGGCTTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box						
6285 CAGGCTTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box						
6285 CAGGCTTTTTG 2,00 0,67 Hs.83484 SRY (sex determining region Y)-box 4 6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box						RNA-binding protein (autoantigenic)
6286 GAAAAGCCTTC 2,00 0,67 Hs.78619 gamma-glutamyl hydrolase (conjugase, folylpolygammag 6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box						SRY (sex determining region Y)-box 4
6287 GTTGAAACTCA 2,00 0,67 Hs.6179 DEAD/H (Asp-Glu-Ala-Asp/His) box	6286	GAAAAGCCTTC	2,00	0,67	Hs.78619	
polypeptide 17 (72k	6287	GTTGAAACTCA	2,00	0,67	Hs.6179	
						polypeptide 17 (72k

6288	GTGCCTAGGAG	2,00	0.67	Hs.25999	Homo sapiens clone 23781 mRNA
					sequence
6289	GGCAACAAAAG	3,00	0,96		karyopherin (importin) beta 1
6290	ATACATTTAGG	1,00	0,34	Hs.77324	eukaryotic translation termination factor 1
6291	CAATTTAAGTG	1,00	0,34	Hs.77324	eukaryotic translation termination factor 1
6292	CTCGCTCCAGG	1,00	0,34		ESTs
6293	ATGGGTCAGAA	1,00	0,34	Hs.66151	Homo sapiens mRNA; cDNA
					DKFZp434A115 (from clone DKF
	AGTGGGTATTT	1,00			CGI-34 protein
6295	GATTGGCGGCT	1,00			actin-like 6
6296	GGAAGTGCAAA	1,00	0,34	Hs.273186	hypothetical protein, clone
222		1.00			Telethon(Italy_B41)_Stra
	GGATAAATGCC	1,00			nucleoporin 153kD
	TGTTTGCATAA	1,00			protein tyrosine kinase 2 beta
	ATTCACCCCCT	1,00			KIAA0561 protein
6300	CTTATAATAAG	1,00	0,34	Hs.129548	heterogeneous nuclear ribonucleoprotein K
6301	CTTTTTCTGT	1,00	0,34	Hs.12013	ATP-binding cassette, sub-family E
	•				(OABP), member 1
	AAGCTCCCTGT	1,00			transducin (beta)-like 3
	CAATCAGAATC	1,00			chromosome 14 open reading frame 2
6304	CTGGCCCCGAG	1,00	0,34	Hs.100997	Human DNA sequence from clone RP1- 18C9 on chromosome
6305	TACGTTGCAGC	3,00	0.06	Hs.21756	translation factor sui1 homolog
	AATGACTGAAT	2,00		Hs.93659	protein disulfide isomerase related
0300	ANIOACIOANI	2,00	0,00	113.33003	protein (calcium
6307	CAGTGTTGGGG	2,00	0.68	Hs.272480	cell matrix adhesion regulator
	GTTTGCAAGTG	2,00			U5 snRNP-specific protein, 116 kD
	GGTGAAGAGGA	3,00			serine protease inhibitor, Kunitz type 1
	TGCTTCATCTG	5,00			RAN, member RAS oncogene family
	CCCTCCTCCGT	2,00		Hs.81131	guanidinoacetate N-methyltransferase
	AGGTGCAGAGG	2,00		Hs.13501	pescadillo (zebrafish) homolog 1,
				·	containing BRCT do
6313	TGCTTTGCTTC	1,00	0,35	Hs.81875	growth factor receptor-bound protein 10
6314	TTTTGTACTTG	1,00		Hs.4245	chromosome 11 hypothetical protein ORF3
6315	AGAAGTACTGA	1,00	0,35	Hs.2934	ribonucleotide reductase M1 polypeptide
6316	GGCGGCTGTGG	1,00			peroxisomal acyl-CoA thioesterase
6317	GCCTGAGGGCC	1,00	0,35	Hs.239500	KIAA0273 gene product
6318	TTGTGTGTACC	1,00	0,35	Hs.208985	ESTs, Weakly similar to plakophilin 2b [H.sapiens]
6319	GAGAGAAAATT	1,00	0.35	Hs.181444	hypothetical protein
	AAATTGTTCCA	1,00			proteasome (prosome, macropain)
					subunit, alpha type,
6321	AACAGCTGGAA	1,00	0,35	Hs.13340	histone acetyltransferase 1
6322	AATAAAAGACC	1,00	0,35	Hs.119178	cation-chloride cotransporter-interacting protein
6323	TCTGCTAAAGA	5,00	1,51	Hs.95958	solute carrier family 2 (facilitated glucose transpo
		1		1	[

6324	GACTCTCTGTT	2,00	0.69	Hs 21635	tubulin, gamma 1
	AAAGGTTGGTT	2,00			ES1 (zebrafish) protein, human homolog
0020	77700110011	2,00	0,03	113.102423	of
6326	GCTTTTAAGGA	17,00	4 31	Hs.8102	ribosomal protein S20
	AAGTCATTCAG	4,00			NADH dehydrogenase (ubiquinone) 1
0027		7,00	1,21	113.274410	alpha subcomplex,
6328	GTGCTTGTACT	2,00	0.70	Hs 151413	glia maturation factor, beta
	AAGGATGCGGT	1,00	0,76	Hs 83126	TATA box binding protein (TBP)-
0025	, 1,00,1,00001	1,00	0,00	110.00120	associated factor, RN
6330	TTTTATGGGTA	1,00	0.36	Hs.79101	cyclin G1
	CCTCCCAGCAA	1,00		Hs.4200	hypothetical protein R32184_1
	CTGTTTAAACT	1,00		Hs.3382	protein phosphatase 4, regulatory
0002	3	,,00	0,00	13.0002	subunit 1
6333	TAGTCATCAAG	1,00	0.36	Hs 281434	sialyltransferase
	GCAATGCAAAA	1,00			KIAA1001 protein
	TCTGCACTGAG	1,00			Homo sapiens mRNA; cDNA
0000		,,00	. 0,00	110.170700	DKFZp434J1313 (from clone DK
6336	GCCGGGCGCGG	1,00	0.36	Hs 155090	hypothetical protein
	CACAAAATCTC	1,00			KIAA0517 protein
	GATTGTGCAAG	2,00			C9orf10 protein
	TTTTGAAGCAG	4,00		Hs.80464	hepatitis B virus x-interacting protein
0000	TTTOARGOAG	7,00	1,20	113.00-10-1	(9.6kD)
6340	GAGCAGCTGGA	2,00	0.72	Hs.166887	
	TTTTTGATAAA	4,00			eukaryotic translation elongation factor 1
0041	TITTOATAVA	7,00	1,50	113.101100	alpha 1
6342	GCCGAGGAAGG	53,00	12 71	He 82148	ribosomal protein S12
	GTTGTCTTTGG	6,00			complement component 3
	AGCAAGTCTCT	1,00			2,4-dienoyl CoA reductase 1,
0011	7.0077.010101	1,00	0,01	113.01040	mitochondrial
6345	CAGTGAATGAA	1,00	0.37	Hs 195614	splicing factor 3b, subunit 3, 130kD
_	CCTGAGGGTAC	1,00			ribosomal protein L8
	CAGCTCTGAGA	1,00			cleavage stimulation factor, 3' pre-RNA,
	0,100,10,10,10,1	1,00	0,01		subunit 1,
6348	TGTAACAATAA	1,00	0.37		Human DNA sequence from clone RP3-
00.0		1,00	0,01	1 13. 12000	322G13 on chromoso
6349	TATCTGTCTAC	5,00	1.58	Hs 145279	SET translocation (myeloid leukemia-
		0,00	.,00		associated)
6350	AAGCTGGAGGA	3,00	1.03		eukaryotic translation initiation factor 3,
		0,00	.,	. ,0.00,002	subunit
6351	GCCTCTGCCAG	2,00	0.72	Hs.106674	BRCA1 associated protein-1 (ubiquitin
		_,	-,		carboxy-termin
6352	ATGAAAAGAAA	3,00	1.04	Hs.76550	Homo sapiens mRNA; cDNA
		.,	.,-		DKFZp564B1264 (from clone DK
6353	TIGITCTTIGT	2,00	0.74	Hs.91773	protein phosphatase 2 (formerly 2A),
		,	- 1.		catalytic subun
6354	TCGTTGTTTAA	1,00	0.37	Hs.8054	CGI-68 protein
	AGGCACTGGCC	1,00		Hs.8008	DNA methyltransferase 1-associated
		','	-,		protein 1
6356	CTGAGTTAGGT	1,00	0,37	Hs.72980	Protein P3
		طسلسا			•

6357	CTCCCGGCGAT	1,00	0,37	Hs.72289	hypothetical protein FLJ20327
6358	TCAGATCCGTC	1,00			Ras-related GTP-binding protein
6359	GGGCGAGAACA	1,00			tumor necrosis factor alpha-inducible
			•		cellular prote
6360	AGGAATGTTAA	1,00	0,37	Hs.250904	zinc finger protein 258
6361	ACATTCCAAGT	1,00	0,37	Hs.245188	tissue inhibitor of metalloproteinase 3
					(Sorsby fund
6362	AATACTTAAAT	1,00	0,37	Hs.241507	ribosomal protein S6
6363	CAGCACATTAT	1,00	0,37	Hs.184592	Human clone A9A2BRB5
<u></u> .					(CAC)n/(GTG)n repeat-containing
6364	CAGAGTGACTG	1,00	0,37	Hs.172589	nuclear phosphoprotein similar to S.
	· · · · · · · · · · · · · · · · · · ·				cerevisiae PWP1
$\overline{}$	GAAATTAGGGA	1,00			KIAA0830 protein
=	TTCTCTCCAAC	1,00			KIAA0250 gene product
	TTCCTCCACCC	1,00		Hs.125743	
	ATTTTCAAAAA	1,00		Hs.102497	
	GCACAGGCCAG	3,00			NEU1 protein
6370	ACCATTCTGCT	3,00	1,06	Hs.174195	interferon induced transmembrane
					protein 2 (1-8D)
	GTTTATGGATA	2,00			matrix Gla protein
	CTGTCATTTGT	3,00			splicing factor, arginine/serine-rich 3
6373	CCTGGAAGAGG	12,00	3,49	Hs.75655	procollagen-proline, 2-oxoglutarate 4-
					dioxygenase (p
6374	CTGTGCTCGGA	3,00	1,07	Hs.76394	enoyl Coenzyme A hydratase, short
	· · · · · · · · · · · · · · · · · · ·			<del></del>	chain, 1, mitochon
6375	GATTAAGTGAG	2,00	0,75	Hs.95835	ESTs, Weakly similar to unnamed
					protein product [H.s
	TGGGCGCCTTT	2,00		Hs.78601	uroporphyrinogen decarboxylase
6377	GAGGTGCÇGGA	2,00	0,75	Hs.37003	v-Ha-ras Harvey rat sarcoma viral
0070	07000070000	0.00			oncogene homolog
	CTGCGGTGGCG	2,00		Hs.252831	
	AAAACATCCAG	1,00		Hs.7763	vesicle docking protein p115
6380	GTTGGCCTGGT	1,00	0,38	Hs.74266	ESTs, Highly similar to similar to
0204	ACCACCTCCAC	4.00	0.00	11- 50007	GTPase-activating
0301	ACGAGCTGGAG	1,00	0,38	Hs.50267	putative GTP-binding protein similar to
6202	CCCACCCTCTC	1.00	0.20	11- 22054	RAY/RAB1C
	GGCAGGCTGTG	1,00			peptidylprolyl isomerase E (cyclophilin E)
0303	GTGTTCCCATA	1,00	0,30		Human DNA sequence from clone RP3-351K20 on chromoso
6384	TTCATTAAAAA	1,00	0.20		Human beta-1D integrin mRNA,
0304	ITCATTAAAAA	1,00	0,36		
6385	GACATTTGTCC	1,00	0.38	Hs.184870	cytoplasmic domain, par
	CGGTTTGCAGA	1,00			Nit protein 2
	GCAAACAATCA	1,00			NS1-associated protein 1
	AACTCCTTCGT	1,00			H2B histone family, member C
	TCTCAAGAAGC				
0008	11010AGAAGC	1,00	0,30	i is. 100000	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 18 (Myc
6390	GCCGTGTCCGC	38,00	10.00	He 2/1507	ribosomal protein S6
	GCCGTGTCCGC	<del></del>			
0001	199010010001	5,00	1,00	Hs.5215_	integrin beta 4 binding protein

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6302	TTCACAGTGGC	5,00	1.67	He 160002	hypothetical 43.2 Kd protein
	TGGCTGTGTGG	4,00		Hs.25709	ESTs, Weakly similar to PSF_HUMAN
0393	16661616166				PTB-ASSOCIATED SPL
	TACAAAAGTGG	2,00			calponin 3, acidic
6395	GGCAGGCGGGT	2,00	0,76	Hs.110906	Ets2 repressor factor
6396	TGGCCCCAGGT	4,00	1,39	Hs.268571	apolipoprotein C-I
6397	ATCAGTGGCTT	6,00	1,97	Hs.89545	proteasome (prosome, macropain)
					subunit, beta type,
6398	GCCTGCAGTCT	7,00			serine protease inhibitor, Kunitz type, 2
6399	GAAAGGTCTGG	3,00	1,10	Hs.118778	KDEL (Lys-Asp-Glu-Leu) endoplasmic
					reticulum protein
6400	AGGAATGCTTT	2,00	0,77	Hs.84131	threonyl-tRNA synthetase
6401	GCTGCTGGTGT	2,00	0,77	Hs.181391	hypothetical protein DKFZp761G2113
6402	TGGAACAGGAT	1,00	0,39	Hs.90077	TG-interacting factor (TALE family
					homeobox)
	CTACACCAGTT	1,00		Hs.71787	30S ribosomal protein S7 homolog
	CAGGATGACGC	1,00			hypothetical protein FLJ10669
6405	AGGGCAGTACT	1,00	0,39	Hs.265561	CD2-associated protein
6406	CCCATCGTCTT	1,00	0,39	Hs.21490	hypothetical protein FLJ20302
6407	CCTCCAGCAGC	1,00	0,39	Hs.17466	retinoic acid receptor responder
	····				(tazarotene induced
6408	GAGTTCGACCT	1,00	0,39	Hs.145362	ESTs, Weakly similar to CHD2_HUMAN CHROMODOMAIN-HELI
6409	CCCGACGTGCC	5,00	1,70	Hs.198269	NADH dehydrogenase (ubiquinone) 1
					alpha subcomplex,
6410	GCCTCTTGAAG	4,00	1,42	Hs.184326	CDC10 (cell division cycle 10, S.
					cerevisiae, homolo
	CACACCCCTGA	2,00			progesterone binding protein
	TAATAAAGCAT	2,00		Hs.4888	seryl-tRNA synthetase
	GGGCAAGCCAG	2,00			estrogen-related receptor alpha
	TGGAAGGGCAC	3,00			EAP30 subunit of ELL complex
6415	CCTCCCCGTC	4,00	1,43	Hs.268763	Breakpoint cluster region protein, uterine leiomyoma
6416	GAGGGGAAACG	2,00	0,78	Hs.81972	SHC (Src homology 2 domain-
					containing) transforming
6417	GGAGGCAGGTG	2,00	0,78	Hs.206713	UDP-Gal:betaGlcNAc beta 1,4-
					galactosyltransferase,
	ACAGACTGATA	1,00			hypothetical protein
	TGGAAAGAGCC	1,00		Hs.9043	DKFZP564O092 protein
	CTTGGTAATTT	1,00		Hs.8768	hypothetical protein FLJ10849
	ACCTGTAATTG	1,00			hqp0256 protein
	TTTCTGTTAAA	1,00			hypothetical protein
6423	GTTGCAGATAA	1,00	0,40	Hs.100293	O-linked N-acetylglucosamine (GlcNAc) transferase (U
6424	AGGTCCTAGCC	9,00	2,87	Hs.226795	glutathione S-transferase pi
6425	CAATGTGTTAT	6,00			NADH dehydrogenase (ubiquinone) 1
			•		alpha subcomplex,
6426	CACAGGCAAAA	4,00	1,44	Hs.155291	KIAA0005 gene product
6427	CTGCACTTACT	4,00		Hs.77152	minichromosome maintenance deficient

6428   CGCATTAAAGC   Co.   C		т —	1		<del></del>	(C. corovinido)
6429 AAGACTGGCTT         2,00         0,79 Hs.284296 Homo sapiens SURF-4 mRNA, complete ods ods           6430 GTGATCTCCGT         2,00         0,79 Hs.158287 KlAA0488 gene product           6431 GCCAAAACCTT         2,00         0,79 Hs.158287 KlAA0488 gene product           6432 TTCTGTGAATC         5,00         1,75 Hs.77870         ESTs           6433 TTACTTCCCCA         3,00         1,13 Hs.184641 deltae fatty acid desaturase           6434 CCCATCCGAAA         22,00         6,47 Hs.91379         ribosomal protein L26           6435 TTCAGGGGAG         6,00         2,06 Hs.9804         DKFZP564C1940 protein           6436 GGCCCCGGACC         4,00         1,46 Hs.4742         anchor attachment protein 1 (Gaa1p, yeast) homolog           6437 CCAAGAAAGAA         2,00         0,80 Hs.169900         poly(A)-binding protein, cytoplasmic 4 (inducible fo           6438 GTGCTGGTGCT         1,00         0,40 Hs.9613         PPAR(gamma) angiopoietin related protein           6439 CCTTTGTAAAA         1,00         0,40 Hs.37189         similar to rat HREV107           6440 TTCTCAAGAAA         1,00         0,40 Hs.3352         histone deacetylase 2           6442 TAGACTTCCTT         1,00         0,40 Hs.3011         ESTs           6444 CATITATCATC         1,00         0,40 Hs.18104         hypothetical protein FLJ11274	6420	CCCATTAAACC	2.00	0.70	Ha 79702	(S. cerevisiae)
Cds		<del></del>				
6431 GCCAAAACCTT         2,00         0,79 Hs.158287 KIAA0468 gene product           6432 ITCGTGAATC         5,00         1,75 Hs.77870 ESTs           6433 ITTACTTCCCA         3,00         1,13 Hs.184641 delta-6 fatty acid desaturase           6434 CCCATCCGAAA         22,00         6,47 Hs.91379 ribosomal protein L26           6435 ITTCAGGGGAG         6,00         2,06 Hs.3804 DKFZP564C1940 protein           6436 GGCCCCGGACC         4,00         1,46 Hs.4742 anchor attachment protein 1 (Gaa1p, yeast) homolog           6437 CCAAGAAAGAA         2,00         0,80 Hs.169900 poly(A)-binding protein, cytoplasmic 4 (inducible fo (inducible fo yeast) homolog           6438 GTGCTGGTGCT         1,00         0,40 Hs.78465 Vjun avian sarcoma virus 17 oncogene homolog           6439 CCTTTGTAAAA         1,00         0,40 Hs.37189 similar to rat HREV107           6440 TTCTCAAGAAA         1,00         0,40 Hs.3352 histone deacetylase 2           6441 CTTTATGTGAT         1,00         0,40 Hs.3352 histone deacetylase 2           6442 TAGACTTCCTT         1,00         0,40 Hs.285999 trinucleotide repeat containing 15           6443 GAGATTTCAATT         1,00         0,40 Hs.18104 hypothetical protein FLJ11274           6444 CATTTATCATC         1,00         0,40 Hs.18104 hypothetical protein FLJ120563           6445 GAGGCTCCTGT         3,00         1,14 Hs.119257 ems1 sequence (mammary tu	0429	AAGACTGGCTT	2,00	0,79	ns.284296	· · · · · · · · · · · · · · · · · · ·
6432 TTCTGTGAATC         5,00         1,75 Hs.77870         ESTs           6433 TTACTTCCCA         3,00         1,13 Hs.184641 delta-6 fatty acid desaturase           6434 CCCATCCGAAA         22,00         6,47 Hs.91379         Pibosomal protein L28           6435 TTTCAGGGGAG         6,00         2,06         Hs.3804         DKFZP564C1940 protein           6436 GGCCCCGGACC         4,00         1,46         Hs.4742         anchor attachment protein 1 (Gaa1p, yeast) homolog           6437 CCAAGAAAGAA         2,00         0,80         Hs.169900 poly(A)-binding protein, cytoplasmic 4 (inducible fo           6438 GTGCTGGTGCT         1,00         0,40         Hs.9813         PPAR(gamma) angiopoietin related protein           6439 CCTTTGTAAAA         1,00         0,40         Hs.78465         Vjun avian sarcoma virus 17 oncogene homolog           6440 TTCTCAAGAAA         1,00         0,40         Hs.3011         ESTs           6442 TAGACTTCCTT         1,00         0,40         Hs.3011         ESTs           6443 JACATTCATT         1,00         0,40         Hs.227777         protein tyrosine phosphatase type IVA, member 1           6445 GATGGCTGCCT         1,00         0,40         Hs.18104         hypothetical protein FLJ11274           6445 GATGCTGCCTG         1,00         0,40			2,00	0,79	Hs.278554	heterochromatin-like protein 1
6433   CCATCCGAAA   22,00   6,47   Hs.91379   ribosomal protein L26   6435   TTACAGGGAG   6,00   2,06   Hs.3804   DKFZP564C1940 protein   Ga15, gest)   CCAAGAAAGAA   2,00   0,80   Hs.16990   Dyly(A)-binding protein, cytoplasmic 4   (Inducible fo   PPAR(gamma) angiopoietin related protein   Ga439   CCTTGTAAAA   1,00   0,40   Hs.78465   V-jun avian sarcoma virus 17 oncogene   homolog   similar to rat HREV107   Hs.3011   ESTs   Ga443   CATTTACATT   1,00   0,40   Hs.285999   trinucleotide repeat containing 15   Ga445   CATTTACATT   1,00   0,40   Hs.285999   trinucleotide repeat containing 15   Ga445   CAGCACTCGTT   1,00   0,40   Hs.18104   Hypothetical protein FLJ11274   Ga447   CTGAAAAAAAAA   1,00   0,40   Hs.18104   Hypothetical protein FLJ20563   Ga449   GAGCTCGCT   1,00   0,40   Hs.18104   Hypothetical protein FLJ20563   Ga450   TCGAAAAAAAA   1,00   0,40   Hs.18104   Hs.19257   Hs.183037   protein kinase, cAMP-dependent, regulatory, type I, Hs.183037   Fortein kinase, cAMP-dependent, regulatory, type I, Hs.183037   Fortein kinase, cAMP-dependent, regulatory, type I, Hs.184062   Putative Rab5-interacting protein 4   Ga450   GAGCACTCCTT   1,00   0,81   Hs.184062   Putative Rab5-interacting protein 4   Ga450   GTGCCAAA   2,00   0,81   Hs.184079   Gytochrome c oxidase subunit VIII   Ga450   GAGCACTCCTT   1,00   0,81   Hs.185073   Gytochrome c oxidase subunit VIII   Ga450   GAGCACTCCCTT   1,00   0,81   Hs.250773   Gytochrome c oxidase subunit VIII   Ga450   GAGCACTCCCAA   2,00   0,81   Hs.250773   Gytochrome c oxidase subunit VIII   Ga450   GAGCACTCCCAA   2,00   0,81   Hs.250773   Gytochrome c oxidase subunit VIII   Ga450   GAGC	6431	GCCAAAACCTT	2,00	0,79	Hs.158287	KIAA0468 gene product
6435 TTTCAGGGAG 6436 GGCCCCGGACC 4,00 1,46 Hs.4742 anchor attachment protein (Gaa1p, yeast) homolog 6437 CCAAGAAAGAA 2,00 0,80 Hs.169900 poly(A)-binding protein, cytoplasmic 4 (inducible fo 6438 GTGCTGGTGCT 1,00 0,40 Hs.9613 6439 CCTTTGTAAAA 1,00 0,40 Hs.78465 6440 TTCTCAAGAAA 1,00 0,40 Hs.37189 6441 CTTTATGTGAT 1,00 0,40 Hs.337189 6442 TAGACTTCCTT 1,00 0,40 Hs.3011 ESTs 6443 ACATTTCAATT 1,00 0,40 Hs.30011 ESTs 6444 CATTTATCATC 1,00 0,40 Hs.385999 trinucleotide repeat containing 15 6445 GATGGCTGCCT 1,00 0,40 Hs.18104 hypothetical protein FLJ11274 6446 TGGAAACAAAA 1,00 0,40 Hs.153834 KIAA0099 gene product 6447 CTGAAAAAAAA 1,00 0,40 Hs.153834 KIAA0099 gene product 6448 GCAGCTCCTGT 3,00 1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci 6450 TGGCTAATAT 3,00 1,15 Hs.183037 Forcin kinase, cAMP-dependent, regulatory, type I, Homo sapiens clone 23704 mRNA sequence 6452 CAGCATCAAT 2,00 0,81 Hs.18029 Hs.18029 Hs.18029 Hs.18029 Hs.18029 Hs.18029 Hs.18030 Friedin kinase, cAMP-dependent, regulatory, type I, Homo sapiens clone 23704 mRNA sequence 6453 GAGACTACAA 2,00 0,81 Hs.18029 Hs.18020	6432	TTCTGTGAATC	5,00	1,75	Hs.77870	ESTs
6435 TTTCAGGGAG 6436 GGCCCCGGACC 4,00 1,46 Hs.4742 anchor attachment protein (Gaa1p, yeast) homolog 6437 CCAAGAAAGAA 2,00 0,80 Hs.169900 poly(A)-binding protein, cytoplasmic 4 (inducible fo 6438 GTGCTGGTGCT 1,00 0,40 Hs.9613 6439 CCTTTGTAAAA 1,00 0,40 Hs.78465 6440 TTCTCAAGAAA 1,00 0,40 Hs.37189 6441 CTTTATGTGAT 1,00 0,40 Hs.337189 6442 TAGACTTCCTT 1,00 0,40 Hs.3011 ESTs 6443 ACATTTCAATT 1,00 0,40 Hs.30011 ESTs 6444 CATTTATCATC 1,00 0,40 Hs.385999 trinucleotide repeat containing 15 6445 GATGGCTGCCT 1,00 0,40 Hs.18104 hypothetical protein FLJ11274 6446 TGGAAACAAAA 1,00 0,40 Hs.153834 KIAA0099 gene product 6447 CTGAAAAAAAA 1,00 0,40 Hs.153834 KIAA0099 gene product 6448 GCAGCTCCTGT 3,00 1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci 6450 TGGCTAATAT 3,00 1,15 Hs.183037 Forcin kinase, cAMP-dependent, regulatory, type I, Homo sapiens clone 23704 mRNA sequence 6452 CAGCATCAAT 2,00 0,81 Hs.18029 Hs.18029 Hs.18029 Hs.18029 Hs.18029 Hs.18029 Hs.18030 Friedin kinase, cAMP-dependent, regulatory, type I, Homo sapiens clone 23704 mRNA sequence 6453 GAGACTACAA 2,00 0,81 Hs.18029 Hs.18020	6433	TTACTTCCCCA	3,00	1,13	Hs.184641	delta-6 fatty acid desaturase
6435 TTTCAGGGAG 6,00 2,06 Hs.3804 DKFZP564C1940 protein anchor attachment protein 1 (Gaa1p, yeast) homolog 6437 CCAAGAAAGAA 2,00 0,80 Hs.169900 poly(A)-binding protein, cytoplasmic 4 (Inducible fo PPAR(gamma) angiopoietin related protein 6438 GTGCTGGTGCT 1,00 0,40 Hs.9613 PPAR(gamma) angiopoietin related protein 6439 CCTTTGTAAAA 1,00 0,40 Hs.37189 similar to rat HREV107 6440 TTCTCAAGAAA 1,00 0,40 Hs.37189 similar to rat HREV107 6441 CTTTATGTGAT 1,00 0,40 Hs.3352 histone deacetylase 2 6442 TAGACTTCCTT 1,00 0,40 Hs.285999 trinucleotide repeat containing 15 6444 CATTTATCATC 1,00 0,40 Hs.227777 protein tyrosine phosphatase type IVA, member 1 6445 GATGGCTGCCT 1,00 0,40 Hs.18104 hypothetical protein FLJ11274 6446 TGGAAACATT C 1,00 0,40 Hs.12142 hypothetical protein FLJ20563 6449 GATGCTGCCAA 12,00 3,78 Hs.199914 ribosomal protein L22 6450 TGTGCTAATAT 3,00 1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type 1, Homo sapiens clone 23704 mRNA sequence 6451 TCCTGAAATAA 2,00 0,81 Hs.18037 protein kinase, cAMP-dependent, regulatory, type 1, Homo sapiens clone 23704 mRNA sequence 6452 CAGCATCTAAT 2,00 0,81 Hs.18003 retinoblastoma-binding protein 4 6453 GAGAGTACAG 2,00 0,81 Hs.18097 cytochrome c oxidase subunit VIII 6456 GTTTTTGCTC 1,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII 6458 GAGCATCCTCTT 1,00 1,48 Hs.79110 nucleolin 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 2,00 0,81 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain-binding protein SH3-domain-binding	6434	CCCATCCGAAA		6,47	Hs.91379	ribosomal protein L26
6436 GGCCCGGACC	6435	TTTCAGGGGAG				
6437 CCAAGAAAGAA 2,00 0,80 Hs.169900 poly(A)-binding protein, cytoplasmic 4 (inducible fo PPAR(gamma) angiopoietin related protein PPAR(gamma) angiopoietin related protein (inducible fo PPAR(gamma) angiopoietin related protein PPAR(gamma) angiopoietin related protein (inducible fo PPAR(gamma) angiopoietin related protein (inducible fo PPAR(gamma) angiopoietin related protein PLJ 100 0,40 Hs.37189 similar to rat HREV107 histone deacetylase 2 (inducible for inducible for indu	6436	GGCCCCGGACC		1,46	Hs.4742	
6438         GTGCTGGTGCT         1,00         0,40 Hs.9613         PPAR(gamma) angiopoietin related protein           6439         CCTTTGTAAAA         1,00         0,40 Hs.78465         V-jun avian sarcoma virus 17 oncogene homolog           6440         TTCTCAAGAAA         1,00         0,40 Hs.37189         similar to rat HREV107           6441         CTTTATGTGAT         1,00         0,40 Hs.3352         histone deacetylase 2           6442         TAGACTTCCTT         1,00         0,40 Hs.285999 trinucleotide repeat containing 15           6443         ACATTTCAATT         1,00         0,40 Hs.28599 trinucleotide repeat containing 15           6444         CATTTATCATC         1,00         0,40 Hs.2818104         hypothetical protein FLJ11274           6445         GATGGCTGCCT         1,00         0,40 Hs.18104         hypothetical protein FLJ20563           6446         TGGAAAAAAAA         1,00         0,40 Hs.19257         ems1 sequence (mammary tumor and squamous cell carci           6449         GATGCTGCCAA         12,00         3,78 Hs.99914         ribosomal protein L22           6450         TGTGCTAATAT         3,00         1,15 Hs.183037         protein kinase, cAMP-dependent, regulatory, type I,           6451         TCCTGAAATAA         2,00         0,81 Hs.18003         retinoblastoma-bindin	6437	CCAAGAAAGAA	2,00	0,80	Hs.169900	poly(A)-binding protein, cytoplasmic 4
6439 CCTTTGTAAAA         1,00         0,40 Hs.78465 v-jun avian sarcoma virus 17 oncogene homolog           6440 TTCTCAAGAAA         1,00         0,40 Hs.37189 similar to rat HREV107           6441 CTTTATGTGAT         1,00         0,40 Hs.3352 histone deacetylase 2           6442 TAGACTTCCTT         1,00         0,40 Hs.30011 ESTs           6443 ACATTTCAATT         1,00         0,40 Hs.285999 trinucleotide repeat containing 15           6444 CATTTATCATC         1,00         0,40 Hs.227777 protein tyrosine phosphatase type IVA, member 1           6445 GATGGCTGCCT         1,00         0,40 Hs.18104 hypothetical protein FLJ11274           6446 TGGAAACTTTC         1,00         0,40 Hs.153834 kIAA0099 gene product           6447 CTGAAAAAAAA         1,00         0,40 Hs.12142 hypothetical protein FLJ20563           6448 GCAGCTCCTGT         3,00         1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci ribosomal protein L22           6450 TGTGCTAATAT         3,00         1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I,           6451 TCCTGAAATAA         2,00         0,81 Hs.5636 Homosapiens clone 23704 mRNA sequence           6452 CAGCATCTAAT         2,00         0,81 Hs.16003 retinoblastoma-binding protein 4           6453 GAGGAGAACAG 2,00         0,81 Hs.16003 retinoblastoma-binding protein 4           6455 GTGACCTCCTT         11,00         3,56	6438	GTGCTGGTGCT	1,00	0,40	Hs.9613	PPAR(gamma) angiopoietin related
6440 TTCTCAAGAAA         1,00         0,40 Hs.37189         similar to rat HREV107           6441 CTTTATGTGAT         1,00         0,40 Hs.3352         histone deacetylase 2           6442 TAGACTTCCTT         1,00         0,40 Hs.285999         trinucleotide repeat containing 15           6444 CATTTATCATC         1,00         0,40 Hs.227777         protein tyrosine phosphatase type IVA, member 1           6445 GATGGCTGCCT         1,00         0,40 Hs.18104         hypothetical protein FLJ11274           6446 TGGAAACTTTC         1,00         0,40 Hs.153834         KIAA0099 gene product           6447 CTGAAAAAAAA         1,00         0,40 Hs.19257         sequence (mammary tumor and squamous cell carci ribosomal protein FLJ20563           6448 GCAGCTCCTGT         3,00         1,14 Hs.119257         sequence (mammary tumor and squamous cell carci ribosomal protein L22           6450 TGTGCTAATAT         3,00         1,15 Hs.183037         protein kinase, cAMP-dependent, regulatory, type I,           6451 TCCTGAAATAA         2,00         0,81 Hs.16003         protein kinase, camp-dependent, regulatory, type I,           6452 CAGCATCTAAT         2,00         0,81 Hs.16003         protein kinase, camp-dependent, regulatory, type I,           6453 GAGGTAACAG         2,00         0,81 Hs.16003         protein kinase, camp-dependent, regulatory, type I,           6455 GTGACC	6439	CCTTTGTAAAA	1,00	0,40		v-jun avian sarcoma virus 17 oncogene
6441 CTTTATGTGAT         1,00         0,40 Hs.3352         histone deacetylase 2           6442 TAGACTTCCTT         1,00         0,40 Hs.30011         ESTs           6443 ACATTTCAATT         1,00         0,40 Hs.285999 trinucleotide repeat containing 15           6444 CATTTATCATC         1,00         0,40 Hs.227777 protein tyrosine phosphatase type IVA, member 1           6445 GATGGCTGCCT         1,00         0,40 Hs.18104 hypothetical protein FLJ11274           6446 TGGAAACTTTC         1,00         0,40 Hs.153834 kIAA0099 gene product           6447 CTGAAAAAAAA         1,00         0,40 Hs.12142 hypothetical protein FLJ20563           6448 GCAGCTCCTGT         3,00         1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci           6449 GATGCTGCCAA         12,00         3,78 Hs.99914 ribosomal protein L22           6450 TGTGCTAATAT         3,00         1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I,           6451 TCCTGAAATAA         2,00         0,81 Hs.184062 putative Rab5-interacting protein           6452 CAGCATCTAAT         2,00         0,81 Hs.184062 putative Rab5-interacting protein           6453 GAGATAACAG         2,00         0,81 Hs.145279 SET translocation (myeloid leukemia-associated)           6455 GTGACCTCCTT         11,00         3,56 Hs.81097 cytochrome c oxidase subunit VIII nucleonin           6456 GTTTTGCTTC <td>6440</td> <td>TTCTCAAGAAA</td> <td>1,00</td> <td>0,40</td> <td></td> <td></td>	6440	TTCTCAAGAAA	1,00	0,40		
6442 TAGACTTCCTT         1,00         0,40 Hs.30011         ESTs           6443 ACATTTCAATT         1,00         0,40 Hs.285999 trinucleotide repeat containing 15           6444 CATTTATCATC         1,00         0,40 Hs.287777 protein tyrosine phosphatase type IVA, member 1           6445 GATGGCTGCCT         1,00         0,40 Hs.18104 hypothetical protein FLJ11274           6446 TGGAAACTTTC         1,00         0,40 Hs.181334 KIAA0099 gene product           6447 CTGAAAAAAAA         1,00         0,40 Hs.12142 hypothetical protein FLJ20563           6448 GCAGCTCCTGT         3,00         1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci           6449 GATGCTGCCAA         12,00         3,78 Hs.99914 ribosomal protein L22           6450 TGTGCTAATAT         3,00         1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I,           6451 TCCTGAAATAA         2,00         0,81 Hs.5636 Homo sapiens clone 23704 mRNA sequence           6452 CAGCATCTAAT         2,00         0,81 Hs.184062 putative Rab5-interacting protein           6453 GAGAGTAACAG         2,00         0,81 Hs.16003 retinoblastoma-binding protein 4           6454 TTTAATTGTGA         2,00         0,81 Hs.145279 SET translocation (myeloid leukemia-associated)           6455 GTGACCTCCTT         11,00         3,56 Hs.81097 cytochrome c oxidase subunit VIII           6456 GTTTTTGCTTC						<u> </u>
6443 ACATTTCAATT 1,00 0,40 Hs.285999 trinucleotide repeat containing 15 6444 CATTTATCATC 1,00 0,40 Hs.227777 protein tyrosine phosphatase type IVA, member 1 6445 GATGGCTGCCT 1,00 0,40 Hs.18104 hypothetical protein FLJ11274 6446 TGGAAAACATTC 1,00 0,40 Hs.153834 KIAA0099 gene product 6447 CTGAAAAAAAA 1,00 0,40 Hs.12142 hypothetical protein FLJ20563 6448 GCAGCTCCTGT 3,00 1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci 6449 GATGCTGCCAA 12,00 3,78 Hs.99914 ribosomal protein L22 6450 TGTGCTAATAT 3,00 1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I, Homo sapiens clone 23704 mRNA sequence 6452 CAGCATCTAAT 2,00 0,81 Hs.184062 putative Rab5-interacting protein 6453 GAGAGTAACAG 2,00 0,81 Hs.184062 putative Rab5-interacting protein 4 6454 TTTAATTGTGA 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4 6455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII 6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin 6457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro						
64444 CATTTATCATC         1,00         0,40 Hs.227777         protein tyrosine phosphatase type IVA, member 1           64445 GATGGCTGCCT         1,00         0,40 Hs.18104         hypothetical protein FLJ11274           6446 TGGAAACATTC         1,00         0,40 Hs.153834         KIAA0099 gene product           6447 CTGAAAAAAAA         1,00         0,40 Hs.12142         hypothetical protein FLJ20563           6448 GCAGCTCCTGT         3,00         1,14 Hs.119257         ems1 sequence (mammary tumor and squamous cell carci           6449 GATGCTGCCAA         12,00         3,78 Hs.99914         ribosomal protein L22           6450 TGTGCTAATAT         3,00         1,15 Hs.183037         protein kinase, cAMP-dependent, regulatory, type I,           6451 TCCTGAAATAA         2,00         0,81 Hs.5636         Homo sapiens clone 23704 mRNA sequence           6452 CAGCATCTAAT         2,00         0,81 Hs.184062         putative Rab5-interacting protein           6453 GAGAGTAACAG         2,00         0,81 Hs.16003         retinoblastoma-binding protein 4           6454 TTTAATTGTGA         2,00         0,81 Hs.145279         SET translocation (myeloid leukemia-associated)           6455 GTGACCTCCTT         11,00         3,56 Hs.81097         cytochrome c oxidase subunit VIII           6456 GTTTTTGCTTC         4,00         1,48 Hs.79110         nuclea						
6445         GATGGCTGCCT         1,00         0,40         Hs.18104         hypothetical protein FLJ11274           6446         TGGAAACTTC         1,00         0,40         Hs.153834         KIAA0099 gene product           6447         CTGAAAAAAAA         1,00         0,40         Hs.12142         hypothetical protein FLJ20563           6448         GCAGCTCCTGT         3,00         1,14         Hs.119257         ems1 sequence (mammary tumor and squamous cell carci           6449         GATGCTGCCAA         12,00         3,78         Hs.99914         ribosomal protein L22           6450         TGTGCTAATAT         3,00         1,15         Hs.183037         protein kinase, cAMP-dependent, regulatory, type I,           6451         TCCTGAAATAA         2,00         0,81         Hs.5636         Homo sapiens clone 23704 mRNA sequence           6452         CAGCATCTAAT         2,00         0,81         Hs.184062         putative Rab5-interacting protein           6453         GAGAGTAACAG         2,00         0,81         Hs.16003         retinoblastoma-binding protein 4           6454         TTTAATTGTGA         2,00         0,81         Hs.145279         SET translocation (myeloid leukemia-associated)           6455         GTGACCTCCTT         11,00         3,56						protein tyrosine phosphatase type IVA,
6446 TGGAAACTTTC         1,00         0,40 Hs.153834 KIAA0099 gene product           6447 CTGAAAAAAAA         1,00         0,40 Hs.12142 hypothetical protein FLJ20563           6448 GCAGCTCCTGT         3,00         1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci           6449 GATGCTGCCAA         12,00         3,78 Hs.99914 ribosomal protein L22           6450 TGTGCTAATAT         3,00         1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I,           6451 TCCTGAAATAA         2,00         0,81 Hs.5636 homo sapiens clone 23704 mRNA sequence           6452 CAGCATCTAAT         2,00         0,81 Hs.184062 putative Rab5-interacting protein           6453 GAGAGTAACAG         2,00         0,81 Hs.16003 retinoblastoma-binding protein 4           6454 TTTAATTGTGA         2,00         0,81 Hs.145279 SET translocation (myeloid leukemia-associated)           6455 GTGACCTCCTT         11,00         3,56 Hs.81097 cytochrome c oxidase subunit VIII           6456 GTTTTTGCTTC         4,00         1,48 Hs.79110 nucleolin           6457 AGGCATTGAAA         3,00         1,16 Hs.151734 nuclear transport factor 2 (placental protein 15)           6458 GAATGAGGACA         3,00         1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain           6459 GATCTCGCAAA         2,00         0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat           6	6445	GATGGCTGCCT	1.00	0.40	Hs.18104	
6447 CTGAAAAAAA 1,00 0,40 Hs.12142 hypothetical protein FLJ20563 6448 GCAGCTCCTGT 3,00 1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci 6449 GATGCTGCCAA 12,00 3,78 Hs.99914 ribosomal protein L22 6450 TGTGCTAATAT 3,00 1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I, 6451 TCCTGAAATAA 2,00 0,81 Hs.5636 Homo sapiens clone 23704 mRNA sequence 6452 CAGCATCTAAT 2,00 0,81 Hs.184062 putative Rab5-interacting protein 6453 GAGAGTAACAG 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4 6454 TTTAATTGTGA 2,00 0,81 Hs.145279 SET translocation (myeloid leukemia-associated) 6455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII nucleolin 6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin nucleolin nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain 6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro						
GAGGTCCTGT 3,00 1,14 Hs.119257 ems1 sequence (mammary tumor and squamous cell carci foliosomal protein L22 fibosomal protein L22 fibosomal protein L22 fibosomal protein kinase, cAMP-dependent, regulatory, type I, fibosomal protein kinase, camped pr						
GATGCTGCCAA 12,00 3,78 Hs.99914 ribosomal protein L22 G450 TGTGCTAATAT 3,00 1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I, G451 TCCTGAAATAA 2,00 0,81 Hs.5636 Homo sapiens clone 23704 mRNA sequence G452 CAGCATCTAAT 2,00 0,81 Hs.184062 putative Rab5-interacting protein G453 GAGAGTAACAG 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4 G454 TTTAATTGTGA 2,00 0,81 Hs.145279 SET translocation (myeloid leukemia-associated) G455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII G456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin G457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) G458 GAATGAGGACA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat G460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro				1,14	Hs.119257	ems1 sequence (mammary tumor and
TGTGCTAATAT 3,00 1,15 Hs.183037 protein kinase, cAMP-dependent, regulatory, type I,  6451 TCCTGAAATAA 2,00 0,81 Hs.5636 Homo sapiens clone 23704 mRNA sequence  6452 CAGCATCTAAT 2,00 0,81 Hs.184062 putative Rab5-interacting protein  6453 GAGAGTAACAG 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4  6454 TTTAATTGTGA 2,00 0,81 Hs.145279 SET translocation (myeloid leukemia-associated)  6455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII  6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin  6457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15)  6458 GAATGAGGACA 2,00 0,81 Hs.250773 reticulocalbin 1, EF-hand calcium binding domain  6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat  6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6449	GATGCTGCCAA	12 00	3 78	Hs 99914	
6451 TCCTGAAATAA  2,00 0,81 Hs.5636 Homo sapiens clone 23704 mRNA sequence 6452 CAGCATCTAAT 2,00 0,81 Hs.184062 putative Rab5-interacting protein 6453 GAGAGTAACAG 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4 6454 TTTAATTGTGA 2,00 0,81 Hs.145279 SET translocation (myeloid leukemia-associated) 6455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII 6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin 6457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain 6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro				1,15	Hs.183037	protein kinase, cAMP-dependent,
6452 CAGCATCTAAT 2,00 0,81 Hs.184062 putative Rab5-interacting protein 6453 GAGAGTAACAG 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4 6454 TTTAATTGTGA 2,00 0,81 Hs.145279 SET translocation (myeloid leukemia-associated) 6455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII 6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin 6457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain 6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6451	TCCTGAAATAA	2,00	0,81	Hs.5636	Homo sapiens clone 23704 mRNA
G453 GAGAGTAACAG 2,00 0,81 Hs.16003 retinoblastoma-binding protein 4 G454 TTTAATTGTGA 2,00 0,81 Hs.145279 SET translocation (myeloid leukemia-associated) G455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII G456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin G457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) G458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain G459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat G460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6452	CAGCATCTAAT	2,00	0,81	Hs.184062	
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6455 GTGACCTCCTT 11,00 3,56 Hs.81097 cytochrome c oxidase subunit VIII 6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin 6457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain 6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6454	TTTAATTGTGA	2,00	0,81	Hs.145279	SET translocation (myeloid leukemia-
6456 GTTTTTGCTTC 4,00 1,48 Hs.79110 nucleolin 6457 AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15) 6458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain 6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6455	GTGACCTCCTT	11,00	3,56	Hs.81097	
AGGCATTGAAA 3,00 1,16 Hs.151734 nuclear transport factor 2 (placental protein 15)  6458 GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain  6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat domain-binding protein SH3-domain-binding pro						
GAATGAGGACA 3,00 1,16 Hs.167791 reticulocalbin 1, EF-hand calcium binding domain  6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat  6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6457	AGGCATTGAAA				nuclear transport factor 2 (placental
6459 GATCTCGCAAA 2,00 0,81 Hs.250773 signal sequence receptor, alpha (translocon-associat 6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6458	GAATGAGGACA	3,00	1,16	Hs.167791	reticulocalbin 1, EF-hand calcium binding
6460 CAGTGGGTGTT 2,00 0,81 Hs.220689 Ras-GTPase-activating protein SH3-domain-binding pro	6459	GATCTCGCAAA	2,00	0,81	Hs.250773	signal sequence receptor, alpha
	6460	CAGTGGGTGTT	2,00	0,81	Hs.220689	Ras-GTPase-activating protein SH3-
	6461	TGGACACAAGC	2,00	0,81	Hs.180832	

0400	TOTO A TO A O A A	4 001	0.44	11. 0004	· · · · · · · · · · · · · · · · · · ·
6462	TGTGATCACAA	1,00	0,41	Hs.9661	proteasome (prosome, macropain) subunit, beta type,
6463	тетестетест	1,00	0.41	Hs.772	glycogen synthase 1 (muscle)
	GACAATGAGAA	1,00		Hs.75253	isocitrate dehydrogenase 3 (NAD+)
0707	ONOM TONOM	1,00	0,41	113.70200	gamma
6465	TTCCCTGCAAG	1,00	0.41	Hs.61628	calcium binding atopy-related
		.,,,,,	<b>0,</b>		autoantigen 1
6466	AAACAAATCAC	1,00	0,41	Hs.173714	MORF-related gene X
	GTTCACTGCAG	1,00			intercellular adhesion molecule 1
					(CD54), human rhin
6468	TTACACTAATA	1,00	0,41	Hs.159	tumor necrosis factor receptor
					superfamily, member 1
	GTAGGAAAGCT	1,00			aminopeptidase puromycin sensitive
	GCGACGAGGCG	19,00		Hs.2017	ribosomal protein L38
	AAGTGATTCTG	3,00	1,19	Hs.180677	zinc finger protein 162
$\overline{}$	GGCCTTTTTT	4,00			H1 histone family, member X
6473	AATATTGAGAA	4,00	1,52	Hs.106673	eukaryotic translation initiation factor 3,
					subunit
	TCATTGTAATG	2,00			GTT1 protein
6475	GTGGCCACGGC	2,00	0,83	Hs.112405	S100 calcium-binding protein A9
2.50					(calgranulin B)
6476	AGACAGAGTGG	2,00	0,83	Hs.103833	ESTs, Weakly similar to AF151869_1
0.477	ATCO 4 4 6 4 6 4 6	1 00	0.40	11 70504	CGI-111 protein [
	ATGCAAGAGAG	1,00		Hs.78521	ESTs
$\overline{}$	GGAGGAATTCA	1,00			cathepsin L
	GCGGGAGCGGG	1,00			KIAA0224 gene product
0480	AGGATTAAAAA	1,00	0,42	Hs.72531	ESTs, Weakly similar to unknown
6481	AAGAGCTAATG	1,00	0.42	Hs.55921	[D.melanogaster]
	GGAAGAGAAGG	1,00			glutamyl-prolyl-tRNA synthetase similar to S. cerevisiae RER1
	GGCTCCTTGAG	1,00			exosome component Rrp46
	TGCCTCGTGAA	1,00	0,42		ESTs
	AGAAGGCCTTG	1,00			mannosidase, alpha, class 1A, member
0400	AGAAGGCC11G	1,00	. 0,42	NS.235114	mannosidase, aipha, class 1A, member
6486	CTTGGGAGGCG	1,00	0.42	Hs 226223	KIAA0618 gene product
	CCCGGTGTGTG	1,00			ESTs, Weakly similar to predicted using
		.,,••	٠, ٠ــ		Genefinder (
6488	CCCGGCTCTTA	1,00	0,42	Hs.184544	
6489	GGTACCCATTT	1,00	0,42	Hs.180383	dual specificity phosphatase 6
6490	GGTACAAATAA	1,00			cAMP responsive element binding
					protein 3 (luman)
6491	CTTCTGCAAAT	1,00	0,42	Hs.14846	Homo sapiens mRNA; cDNA
ا ن					DKFZp564D016 (from clone DKF
	ACCCAGCGGGC	1,00		Hs.126705	ESTs
	GGAGCACTGTG	1,00	0,42	Hs.10362	ESTs
	GAGAATTAATC	1,00			basic transcription factor 3
6495	GAAAAAAAAA	12,00	3,99	Hs.14839	polymerase (RNA) II (DNA directed)
2425	070000				polypeptide G
6496	GTGGGTGTCCT	2,00	0,84	Hs.27633	DKFZP586B0519 protein

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acytransferase 1 (lyso   6498 GCCAAGCCTGA   2,00   0,84   Hs. 118796   annexin A6   6498 AAAGGAAAGTC   2,00   0,84   Hs. 109706   HN1 protein   6500   GGGGTAAGAAA   8,00   2,81   Hs. 80423   prostatic binding protein   6501   TGTCATCACAG   2,00   0,84   Hs. 83354   hypothetical protein DKFZp762l166   6503   GACTCTGGGAT   2,00   0,84   Hs. 8191   hypothetical protein DKFZp762l166   6504   AAAGGGGGCAG   3,00   1,21   Hs. 249247   heterogeneous nuclear protein similar to rat helix d   heterogeneous nuclear ribonucleoprotein   A2/B1   heterogeneous nuclear ribonucleoprotein   heterogeneous	6497	ACCCCACCCAG	2,00	0.84	Hs 240534	1-acylglycerol-3-phosphate O-
6498 GCCAAGCCTGA         2,00         0,84 Hs. 118796 annexin A6           6499 AAAGGAAAGTC         2,00         0,84 Hs. 109706 HN1 protein           6500 GGGGTAAGAAA         8,00         2,31 Hs.80423 prostatic binding protein           6501 TGTCATCACAG         2,00         0,84 Hs.83354 prostatic binding protein DKFZp762l166           6502 GAGAACGGGGA         2,00         0,84 Hs.6191 protein DKFZp762l166           6503 GACTCTGGGAT         2,00         0,84 Hs.11282 ESTs, Weakly similar to cleft lip and palate transme           6504 AAAGGGGCAG         3,00         1,21 Hs.249247 heterogeneous nuclear ribonucleoprotein rat helix d           6505 GTGTAATAAGA         7,00         2,52 Hs.75598 heterogeneous nuclear ribonucleoprotein A2/B1           6506 TGTATTACAGT         1,00         0,43 Hs.78948 Rab geranylgeranyltransferase, beta subunit           6507 CCTGAAGAAGT         1,00         0,43 Hs.74579 KIAA0283 gene product           6508 CACGTTCCCTA         1,00         0,43 Hs.74579 KIAA0283 gene product           6509 AGGGATGGCC         1,00         0,43 Hs.279518 amyloid precursor protein homolog HSD-2           6510 CTCTCAATATA         1,00         0,43 Hs.179574 protein phosphatase 2 (formerly 2A), regulatory subu           6511 TAAACTTTGTA         1,00         0,43 Hs.173374 Homo sapiens cDNA FLJ10500 fis, clone NT2RP-2000369           6512 CTGGAGGCTGC         1,00 <td>0.0.</td> <td>, 10000, 1000, 10</td> <td>2,00</td> <td>0,0 1</td> <td>10.2-1000</td> <td></td>	0.0.	, 10000, 1000, 10	2,00	0,0 1	10.2-1000	
6399 AAAGGAAAGTC         2,00         0,84 Hs. 309706 HN1 protein           6500 GGGTAAGAAA         8,00         2,81 Hs. 80423         prostatic binding protein           6501 TGTCATCACAG         2,00         0,84 Hs. 83354         lysyl oxidase-like 2           6502 GAGAACGGGA         2,00         0,84 Hs. 6191         hypothetical protein DKFZp762I166           6503 GACTCTGGGAT         2,00         0,84 Hs. 6191         hypothetical protein DKFZp762I166           6504 AAAGGGGGCAG         3,00         1,21 Hs. 249247         heterogeneous nuclear ribonucleoprotein palate transme           6505 GTGTAATAAGA         7,00         2,52 Hs. 75598         heterogeneous nuclear ribonucleoprotein A2/B1           6506 TGTATTACAGT         1,00         0,43 Hs. 78948         Rab geranylgeranyltransferase, beta subunit           6507 CCTGAAGAAGT         1,00         0,43 Hs. 74579         KIAA0283 gene product           6508 CACGTTCCCTA         1,00         0,43 Hs. 74579         KIAA0283 gene product           6509 AGGGATGGCCC         1,00         0,43 Hs. 227891         ESTS, Weakly similar to Y167_HUMAN           6511 TAAACTTTGTA         1,00         0,43 Hs. 179574         protein phosphatase 2 (formerly 2A), regulatory subu           6512 CTATGGATTT         1,00         0,43 Hs. 179574         protein phosphatase c (formerly 2A), regulatory subu	6498	GCCAAGCCTGA	2.00	0.84	Hs.118796	
BSD0   GGGGTAAGAAA   8,00   2,81   Hs.80423   prostatic binding protein   3501   TGTCATCACAG   2,00   0,84   Hs.81354   lysyl oxidase-like 2   6502   GAGAACGGGGA   2,00   0,84   Hs.81354   lysyl oxidase-like 2   6503   GACTCTGGGAT   2,00   0,84   Hs.11282   ESTs, Weakly similar to cleft lip and palate transme   6504   AAAGGGGGCAG   3,00   1,21   Hs.249247   heterogeneous nuclear protein similar to rat helix d   6505   GTGTAATAAGA   7,00   2,52   Hs.75598   heterogeneous nuclear ribonucleoprotein   A2/81   heterogeneous nuclear ribonucleoprotein   heterogeneous nuclear ribonucleoprotein   A2/81   heterogeneous nuclear ribonucleoprotein   heterogeneous nuclear		<del></del>				
B6501   TGTCATCACAG   2,00   0,84   Hs.83354   yeyl oxidase-like 2   6502   GAGAACGGGGA   2,00   0,84   Hs.6191   hypothetical protein DKFZp762l168   6503   GACTCTGGGAT   2,00   0,84   Hs.11282   ESTs, Weakly similar to cleft lip and palate transme   6504   AAAGGGGGCAG   3,00   1,21   Hs.249247   heterogeneous nuclear protein similar to rat helix d   heterogeneous nuclear ribonucleoprotein Az/B1   6505   GTGTAATAAGA   7,00   2,52   Hs.75598   heterogeneous nuclear ribonucleoprotein Az/B1   6506   TGTATTACAGT   1,00   0,43   Hs.78948   Rab geranylgeranyltransferase, beta subunit   6508   CACGTTCCCTA   1,00   0,43   Hs.74579   KIAAO263 gene product   6509   AGGGATGGCC   1,00   0,43   Hs.279518   amyloid precursor protein homolog HSD-2   2   6511   TAAACTTTGTA   1,00   0,43   Hs.279518   amyloid precursor protein homolog HSD-2   2   6512   CTATGGGATT   1,00   0,43   Hs.179574   protein phosphatase 2 (formerly 2A), regulatory subu   6513   TACTGTAGTCA   1,00   0,43   Hs.179374   homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369   6514   CTGGAGGCTGC   1,00   0,43   Hs.179374   homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369   6515   GAGTCTGAGGG   2,00   0,85   Hs.174051   small nuclear ribonucleoprotein 70kD polypeptide (RN 6519   TGTAATGTTA   2,00   0,86   Hs.244287   putative integral membrane transporter   6519   TGTAATGTTA   2,00   0,43   Hs.248287   putative integral membrane transporter   6521   GGCAGCCTGGT   1,00   0,43   Hs.26438   Bottom sapiens mRNA; cDNA   DKFZp434N0211 (from clone DK   6521   GGCAGCCTGGT   1,00   0,43   Hs.26438   GST   NADH dehydrogenase (ubiquinone)   flavoprotein 3 (10kD   6525   GCTGGAACGAA   1,00   0,43   Hs.26343   EST   6525   GCTGGAACGAA   1,00   0,43   Hs.26343   EST   6526   GCTGGAACGAA   1,00   0,43   Hs.26343   EST   6526   GCTGGACGAC   1,00   0,43   Hs.26343   EST   6526   GCTGGACGAC   1,00   0,43   Hs.26343   EST   6526   GCTGGAACGAC   1,00   0,43   Hs.26343   EST   6526   GCTGGACGAC   1,00   0,43   Hs.26343   EST   6526   GCTGGACGAC   1,00   0,43   Hs.26343   EST   6						
Section						
Section	6502	GAGAACGGGGA		0,84	Hs.6191	hypothetical protein DKFZp762I166
palate transme   palate transmarter   palate   palate transme   palate transme   palate transme   palate transme   palate transme   palate transme   palate transporter   palate   palate						
						palate transme
Section   Continue	6504	AAAGGGGGCAG	3,00	1,21	Hs.249247	heterogeneous nuclear protein similar to
A2/81   A2/8						
1,00	6505	GTGTAATAAGA	7,00	2,52	Hs.75598	
CCTGAAGAAGT   1,00						L
Subunit   Subu						
6508         CACGTTCCCTA         1,00         0,43         Hs.74579         KIAA0263 gene product           6509         AGGGATGGCCC         1,00         0,43         Hs.54411         putative T1/ST2 receptor binding protein           6510         CTCTCAATATA         1,00         0,43         Hs.279518         amyloid precursor protein homolog HSD-2           6511         TAAACTTTGTA         1,00         0,43         Hs.27891         ESTs, Weakly similar to Y167_HUMAN HYPOTHETICAL PROT           6512         CTATGGGATTT         1,00         0,43         Hs.179574         protein phosphatase 2 (formerly 2A), regulatory subu           6513         TACTGTAGTCA         1,00         0,43         Hs.173374         Homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369           6514         CTGGAGGCTGC         1,00         0,43         Hs.174951         ESTs, Weakly similar to rhophilin [M.musculus]           6516         GAGTCTGAGGG         1,00         0,43         Hs.174951         small nuclear ribonucleoprotein 70kD polypeptide (RN           6517         GTGCGCTAGGG         8,00         2,86         Hs.9408         IKK-related kinase epsilon; inducible likappa kinase           6518         GAAAAGGGTTT         2,00         0,86         Hs.284287         putative integral membrane transporter	6507	CCTGAAGAAGT	1,00	0,43	Hs.78948	
6509         AGGGATGGCCC         1,00         0,43 Hs.54411         putative T1/ST2 receptor binding protein           6510         CTCTCAATATA         1,00         0,43 Hs.279518         amyloid precursor protein homolog HSD-2           6511         TAAACTTTGTA         1,00         0,43 Hs.227891         ESTs, Weakly similar to Y167_HUMAN HYPOTHETICAL PROT           6512         CTATGGGATTT         1,00         0,43 Hs.179574         protein phosphatase 2 (formerly 2A), regulatory subu           6513         TACTGTAGTCA         1,00         0,43 Hs.173374         Homo sapiens cDNA FLJ10500 fis, clone. NT2RP2000369           6514         CTGGAGGCTGC         1,00         0,43 Hs.149152         ESTs, Weakly similar to rhophilin [M.musculus]           6515         GCCCCTGAAGG         1,00         0,43 Hs.174051         small nuclear ribonucleoprotein 70kD polypeptide (RN)           6516         GAGTCTGAGGG         8,00         2,86 Hs.9408         IKK-related kinase epsilon; inducible lkappaB kinase           6518         GAAAAGGGTTT         2,00         0,86 Hs.284287 putative integral membrane transporter           6519         TGTTAATGTTA         2,00         0,86 Hs.261828 Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK calcium (ca2+) ho           6520         TTCAGTGCCCA         1,00         0,43 Hs.6430         protein with polyglutamine repeat; calcium (						
6510         CTCTCAATATA         1,00         0,43 Hs.279518 amyloid precursor protein homolog HSD-2           6511         TAAACTTTGTA         1,00         0,43 Hs.227891 ESTs, Weakly similar to Y167_HUMAN HYPOTHETICAL PROT           6512         CTATGGGATTT         1,00         0,43 Hs.179574 protein phosphatase 2 (formerly 2A), regulatory subu           6513         TACTGTAGTCA         1,00         0,43 Hs.173374 Homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369           6514         CTGGAGGCTGC         1,00         0,43 Hs.149152 ESTs, Weakly similar to rhophilin [M.musculus]           6515         GCCCCTGAAGG         1,00         0,43 Hs.174051 small nuclear ribonucleoprotein 70kD polypeptide (RN           6516         GAGTCTGAGGG         2,00         0,85 Hs.174051 small nuclear ribonucleoprotein 70kD polypeptide (RN           6517         GTGCGCTAGGG         8,00         2,86 Hs.9408 lKK-related kinase epsilon; inducible lkappaB kinase           6518         GAAAAGGGTTT         2,00         0,86 Hs.284287 putative integral membrane transporter           6519         TGTTAATGTTA         2,00         0,86 Hs.261828 Homo saplens mRNA; cDNA DKFZp434N0211 (from clone DK cytochrome c oxidase subunit VIc           6520         TTCAGTGCCCA         1,00         0,43 Hs.6430 protein with polyglutamine repeat; calcium (ca2+) ho           6521         GCAGCCTGGT         1,00         0,43 Hs.26						
2						
HYPOTHETICAL PROT	6510	CTCTCAATATA	1,00	0,43	Hs.279518	amyloid precursor protein homolog HSD-
HYPOTHETICAL PROT						2
6512         CTATGGGATTT         1,00         0,43         Hs.179574         protein phosphatase 2 (formerly 2A), regulatory subu           6513         TACTGTAGTCA         1,00         0,43         Hs.173374         Homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369           6514         CTGGAGGCTGC         1,00         0,43         Hs.149152         ESTs, Weakly similar to rhophilin [M.musculus]           6515         GCCCCTGAAGG         1,00         0,43         Hs.117682         CGI-43 protein           6516         GAGTCTGAGGG         2,00         0,85         Hs.174051         small nuclear ribonucleoprotein 70kD polypeptide (RN           6517         GTGCGCTAGGG         8,00         2,86         Hs.9408         IKK-related kinase epsilon; inducible lkappaB kinase           6518         GAAAAGGGTTT         2,00         0,86         Hs.284287 putative integral membrane transporter           6519         TGTTAATGTTA         2,00         0,48         Hs.261828         Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK           6520         TTCAGTGCCCA         1,00         0,43         Hs.6430         protein with polyglutamine repeat; calcium (ca2+) ho           6521         GGCAGCTGGT         1,00         0,43         Hs.274317         DEAD-box protein abstrakt           6524	6511	TAAACTTTGTA	1,00	0,43	Hs.227891	
regulatory subu						
TACTGTAGTCA	6512	CTATGGGATTT	1,00	0,43	Hs.179574	
NT2RP2000369						
6514         CTGGAGGCTGC         1,00         0,43         Hs.149152         ESTs, Weakly similar to rhophilin [M.musculus]           6515         GCCCCTGAAGG         1,00         0,43         Hs.117582         CGI-43 protein           6516         GAGTCTGAGGG         2,00         0,85         Hs.174051         small nuclear ribonucleoprotein 70kD polypeptide (RN           6517         GTGCGCTAGGG         8,00         2,86         Hs.9408         IKK-related kinase epsilon; inducible lkappaB kinase           6518         GAAAAGGGTTT         2,00         0,86         Hs.284287 putative integral membrane transporter           6519         TGTTAATGTTA         2,00         0,86         Hs.261828         Homoo sapiens mRNA; cDNA DKFZP434N0211 (from clone DK cytochrome c oxidase subunit VIc           6520         TTCAGTGCCCA         1,00         0,43         Hs.74649         cytochrome c oxidase subunit VIc           6521         GGCAGCCTGGT         1,00         0,43         Hs.6430         protein with polyglutamine repeat; calcium (ca2+) ho           6522         ACTACAGCACG         1,00         0,43         Hs.59745         NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD           6523         GACTTCTGAGC         1,00         0,43         Hs.26343         ESTs           6524         TTGGTGA	6513	TACTGTAGTCA	1,00	0,43	Hs.173374	
[M.musculus]   [M.m		070010000			11 1101-0	
6515         GCCCCTGAAGG         1,00         0,43         Hs.117582         CGI-43 protein           6516         GAGTCTGAGGG         2,00         0,85         Hs.174051         small nuclear ribonucleoprotein 70kD polypeptide (RN           6517         GTGCGCTAGGG         8,00         2,86         Hs.9408         IKK-related kinase epsilon; inducible lkappaB kinase           6518         GAAAAGGGTTT         2,00         0,86         Hs.284287 putative integral membrane transporter           6519         TGTTAATGTTA         2,00         0,86         Hs.261828         Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK           6520         TTCAGTGCCCA         1,00         0,43         Hs.74649         cytochrome c oxidase subunit VIc           6521         GGCAGCCTGGT         1,00         0,43         Hs.6430         protein with polyglutamine repeat; calcium (ca2+) ho           6522         ACTACAGCACG         1,00         0,43         Hs.274317         DEAD-box protein abstrakt           6524         TTGGTGCTTGG         1,00         0,43         Hs.26343         ESTs           6525         ACTGGAACGAA         1,00         0,43         Hs.11223         isocitrate dehydrogenase 1 (NADP+), soluble           6526         GCTTGATGTGA         1,00         0,43 <t< td=""><td>6514</td><td>CTGGAGGCTGC</td><td>1,00</td><td>0,43</td><td>Hs.149152</td><td></td></t<>	6514	CTGGAGGCTGC	1,00	0,43	Hs.149152	
6516 GAGTCTGAGGG 2,00 0,85 Hs.174051 small nuclear ribonucleoprotein 70kD polypeptide (RN 6517 GTGCGCTAGGG 8,00 2,86 Hs.9408 IKK-related kinase epsilon; inducible lkappaB kinase 6518 GAAAAGGGTTT 2,00 0,86 Hs.284287 putative integral membrane transporter 6519 TGTTAATGTTA 2,00 0,86 Hs.261828 Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK 6520 TTCAGTGCCCA 1,00 0,43 Hs.74649 cytochrome c oxidase subunit VIc 6521 GGCAGCCTGGT 1,00 0,43 Hs.6430 protein with polyglutamine repeat; calcium (ca2+) ho 6522 ACTACAGCACG 1,00 0,43 Hs.59745 NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD 6523 GACTTCTGAGC 1,00 0,43 Hs.274317 DEAD-box protein abstrakt 6524 TTGGTGCTTGG 1,00 0,43 Hs.26343 ESTs 6525 ACTGGAACGAA 1,00 0,43 Hs.11223 isocitrate dehydrogenase 1 (NADP+), soluble 6526 GCTTGATGTGA 1,00 0,43 Hs.108548 hypothetical protein 6527 AAGCTGCTGGA 3,00 1,24 Hs.9822 HCNP protein 6528 CTGGGACTGAC 3,00 1,24 Hs.76719 U6 snRNA-associated Sm-like protein	0546	00000704400	4.00	0.40	11 447500	<del>                                     </del>
Dolypeptide (RN     CTGCGCTAGGG   S,00   2,86   Hs.9408   IKK-related kinase epsilon; inducible lkappaB kinase						
6517 GTGCGCTAGGG 8,00 2,86 Hs.9408 IKK-related kinase epsilon; inducible lkappaB kinase 6518 GAAAAGGGTTT 2,00 0,86 Hs.284287 putative integral membrane transporter 6519 TGTTAATGTTA 2,00 0,86 Hs.261828 Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK 6520 TTCAGTGCCCA 1,00 0,43 Hs.74649 cytochrome c oxidase subunit VIc 6521 GGCAGCCTGGT 1,00 0,43 Hs.6430 protein with polyglutamine repeat; calcium (ca2+) ho 6522 ACTACAGCACG 1,00 0,43 Hs.59745 NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD 6523 GACTTCTGAGC 1,00 0,43 Hs.274317 DEAD-box protein abstrakt 6524 TTGGTGCTTGG 1,00 0,43 Hs.26343 ESTs 6525 ACTGGAACGAA 1,00 0,43 Hs.11223 isocitrate dehydrogenase 1 (NADP+), soluble 6526 GCTTGATGTGA 1,00 0,43 Hs.108548 hypothetical protein 6527 AAGCTGCTGGA 3,00 1,24 Hs.9822 HCNP protein 6528 CTGGGACTGAC 3,00 1,24 Hs.76719 U6 snRNA-associated Sm-like protein	6516	GAGICIGAGGG	2,00	0,85	Hs.1/4051	
IkappaB kinase   6518 GAAAAGGGTTT   2,00   0,86 Hs.284287 putative integral membrane transporter   6519 TGTTAATGTTA   2,00   0,86 Hs.261828 Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK   6520 TTCAGTGCCA   1,00   0,43 Hs.74649   cytochrome c oxidase subunit VIc   6521 GGCAGCCTGGT   1,00   0,43 Hs.6430   protein with polyglutamine repeat; calcium (ca2+) ho   6522 ACTACAGCACG   1,00   0,43 Hs.59745   NADH dehydrogenase (ubiquinone)   flavoprotein 3 (10kD   6523 GACTTCTGAGC   1,00   0,43 Hs.274317 DEAD-box protein abstrakt   6524 TTGGTGCTTGG   1,00   0,43 Hs.26343   ESTs   6525 ACTGGAACGAA   1,00   0,43 Hs.11223   isocitrate dehydrogenase 1 (NADP+), soluble   6526 GCTTGATGTGA   1,00   0,43 Hs.108548   hypothetical protein   6527 AAGCTGCTGGA   3,00   1,24 Hs.9822   HCNP protein   6528 CTGGGACTGAC   3,00   1,24 Hs.76719   U6 snRNA-associated Sm-like protein	6517	GTGCGCTAGGG	8.00	2.86	He 0408	
6518 GAAAAGGGTTT         2,00         0,86 Hs.284287 putative integral membrane transporter           6519 TGTTAATGTTA         2,00         0,86 Hs.261828 Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK           6520 TTCAGTGCCCA         1,00         0,43 Hs.74649 cytochrome c oxidase subunit VIc           6521 GGCAGCCTGGT         1,00         0,43 Hs.6430 protein with polyglutamine repeat; calcium (ca2+) ho           6522 ACTACAGCACG         1,00         0,43 Hs.59745 NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD           6523 GACTTCTGAGC         1,00         0,43 Hs.274317 DEAD-box protein abstrakt           6524 TTGGTGCTTGG         1,00         0,43 Hs.26343 ESTs           6525 ACTGGAACGAA         1,00         0,43 Hs.11223 isocitrate dehydrogenase 1 (NADP+), soluble           6526 GCTTGATGTGA         1,00         0,43 Hs.108548 hypothetical protein           6527 AAGCTGCTGGA         3,00         1,24 Hs.9822 HCNP protein           6528 CTGGGACTGAC         3,00         1,24 Hs.76719 U6 snRNA-associated Sm-like protein	0317	01000017000	0,00	2,00	113.9400	
6519 TGTTAATGTTA         2,00         0,86 Hs.261828 Homo sapiens mRNA; cDNA DKFZp434N0211 (from clone DK           6520 TTCAGTGCCCA         1,00         0,43 Hs.74649 cytochrome c oxidase subunit VIc           6521 GGCAGCCTGGT         1,00         0,43 Hs.6430 protein with polyglutamine repeat; calcium (ca2+) ho           6522 ACTACAGCACG         1,00         0,43 Hs.59745 NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD           6523 GACTTCTGAGC         1,00         0,43 Hs.274317 DEAD-box protein abstrakt           6524 TTGGTGCTTGG         1,00         0,43 Hs.26343 ESTs           6525 ACTGGAACGAA         1,00         0,43 Hs.11223 isocitrate dehydrogenase 1 (NADP+), soluble           6526 GCTTGATGTGA         1,00         0,43 Hs.108548 hypothetical protein           6527 AAGCTGCTGGA         3,00         1,24 Hs.9822 HCNP protein           6528 CTGGGACTGAC         3,00         1,24 Hs.76719 U6 snRNA-associated Sm-like protein	6518	GAAAAGGGTTT	2.00	0.86	Hs 284287	
DKFZp434N0211 (from clone DK						
6520 TTCAGTGCCCA1,000,43 Hs.74649cytochrome c oxidase subunit VIc6521 GGCAGCCTGGT1,000,43 Hs.6430protein with polyglutamine repeat; calcium (ca2+) ho6522 ACTACAGCACG1,000,43 Hs.59745NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD6523 GACTTCTGAGC1,000,43 Hs.274317 DEAD-box protein abstrakt6524 TTGGTGCTTGG1,000,43 Hs.26343ESTs6525 ACTGGAACGAA1,000,43 Hs.11223isocitrate dehydrogenase 1 (NADP+), soluble6526 GCTTGATGTGA1,000,43 Hs.108548 hypothetical protein6527 AAGCTGCTGGA3,001,24 Hs.9822HCNP protein6528 CTGGGACTGAC3,001,24 Hs.76719U6 snRNA-associated Sm-like protein	00.0		2,00	0,00	113.201020	
6521         GGCAGCCTGGT         1,00         0,43         Hs.6430         protein with polyglutamine repeat; calcium (ca2+) ho           6522         ACTACAGCACG         1,00         0,43         Hs.59745         NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD           6523         GACTTCTGAGC         1,00         0,43         Hs.274317         DEAD-box protein abstrakt           6524         TTGGTGCTTGG         1,00         0,43         Hs.26343         ESTs           6525         ACTGGAACGAA         1,00         0,43         Hs.11223         isocitrate dehydrogenase 1 (NADP+), soluble           6526         GCTTGATGTGA         1,00         0,43         Hs.108548         hypothetical protein           6527         AAGCTGCTGGA         3,00         1,24         Hs.9822         HCNP protein           6528         CTGGGACTGAC         3,00         1,24         Hs.76719         U6 snRNA-associated Sm-like protein	6520	TTCAGTGCCCA	1.00	0.43	Hs 74649	
Calcium (ca2+) ho				0.43	Hs 6430	protein with polyalutamine repeat:
6522         ACTACAGCACG         1,00         0,43         Hs.59745         NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD           6523         GACTTCTGAGC         1,00         0,43         Hs.274317         DEAD-box protein abstrakt           6524         TTGGTGCTTGG         1,00         0,43         Hs.26343         ESTs           6525         ACTGGAACGAA         1,00         0,43         Hs.11223         isocitrate dehydrogenase 1 (NADP+), soluble           6526         GCTTGATGTGA         1,00         0,43         Hs.108548         hypothetical protein           6527         AAGCTGCTGGA         3,00         1,24         Hs.9822         HCNP protein           6528         CTGGGACTGAC         3,00         1,24         Hs.76719         U6 snRNA-associated Sm-like protein	0021	000,10001001	,,00	0, 10	110.0100	
	6522	ACTACAGCACG	1.00	0.43	Hs 59745	
6523 GACTTCTGAGC         1,00         0,43 Hs.274317 DEAD-box protein abstrakt           6524 TTGGTGCTTGG         1,00         0,43 Hs.26343 ESTs           6525 ACTGGAACGAA         1,00         0,43 Hs.11223 isocitrate dehydrogenase 1 (NADP+), soluble           6526 GCTTGATGTGA         1,00         0,43 Hs.108548 hypothetical protein           6527 AAGCTGCTGGA         3,00         1,24 Hs.9822 HCNP protein           6528 CTGGGACTGAC         3,00         1,24 Hs.76719 U6 snRNA-associated Sm-like protein			.,55	٥, .٠	.,0.93, 13	
6524 TTGGTGCTTGG         1,00         0,43 Hs.26343         ESTs           6525 ACTGGAACGAA         1,00         0,43 Hs.11223         isocitrate dehydrogenase 1 (NADP+), soluble           6526 GCTTGATGTGA         1,00         0,43 Hs.108548 hypothetical protein           6527 AAGCTGCTGGA         3,00         1,24 Hs.9822         HCNP protein           6528 CTGGGACTGAC         3,00         1,24 Hs.76719         U6 snRNA-associated Sm-like protein	6523	GACTTCTGAGC	1.00	0.43	Hs.274317	
6525         ACTGGAACGAA         1,00         0,43         Hs.11223         isocitrate dehydrogenase 1 (NADP+), soluble           6526         GCTTGATGTGA         1,00         0,43         Hs.108548         hypothetical protein           6527         AAGCTGCTGGA         3,00         1,24         Hs.9822         HCNP protein           6528         CTGGGACTGAC         3,00         1,24         Hs.76719         U6 snRNA-associated Sm-like protein						
6526         GCTTGATGTGA         1,00         0,43         Hs.108548         hypothetical protein           6527         AAGCTGCTGGA         3,00         1,24         Hs.9822         HCNP protein           6528         CTGGGACTGAC         3,00         1,24         Hs.76719         U6 snRNA-associated Sm-like protein			.,,	-,		
6527 AAGCTGCTGGA 3,00 1,24 Hs.9822 HCNP protein 6528 CTGGGACTGAC 3,00 1,24 Hs.76719 U6 snRNA-associated Sm-like protein	6526	GCTTGATGTGA	1.00	0.43	Hs.108548	<del>                                      </del>
6528 CTGGGACTGAC 3,00 1,24 Hs.76719 U6 snRNA-associated Sm-like protein						<del></del>
			2,00			STIP1 homology and U-Box containing

					Investoria de
0000	040070007	44.00		11 470404	protein 1
$\rightarrow$	GACCCTGCCCT	11,00			FK506-binding protein 8 (38kD)
	GTTCGTGCCAA	20,00			ribosomal protein L35a
6532	GACCAGAAAAA	5,00	1,95		cytochrome c oxidase subunit Vla
0500	OTO 4 4 0 0 0 4 0 T	00.00	40.70		polypeptide 1
	GTGAAGGCAGT	33,00			ribosomal protein S3A
6534	CAGATCTTTGT	9,00	3,29		ubiquitin A-52 residue ribosomal protein
CEOF	TOCOCTOTOAA	4 00	0.44		fusion prod
0535	TGGGCTCTGAA	1,00	0,44		CD36 antigen (collagen type I receptor,
GEOC!	OTTOCOCATOO	1.00	0.44		thrombospond
-	CTTCGCGATGG	1,00			KIAA0081 protein
	GAGGTCCTTCG	1,00			inositol 1,3,4-triphosphate 5/6 kinase
$\rightarrow$	GTACATTGTAA	1,00			ESTs
	GTTAATTGCTA	1,00			AD-015 protein
	GCTTATAAAAA	1,00			histidine triad nucleotide-binding protein
	CATACACACAC	1,00		Hs.254279	
6542	тттстстссст	1,00	0,44		SWI/SNF related, matrix associated,
0540	00044744407	4.00	0.44		actin dependent
	CCCAATAAACT	1,00			amida, partner of the E2A
	TAACAGTTGTG	1,00		Hs.155560	
	CTGCCATAACT	2,00			CGI-97 protein
	TTCTCTCCCCA	2,00			DKFZP566D193 protein
654/	CCCAGGGAGAA	3,00	1,27	Hs.79150	chaperonin containing TCP1, subunit 4 (delta)
6548	TGGCCTGCCCA	3,00	1.27	Hs 181002	MLL septin-like fusion (NOTE: non-
		","	.,		standard symbol an
6549	TTACCTCCTTC	9,00	3.34	Hs.3343	phosphoglycerate dehydrogenase
	GCCAGCCCAGC	7,00			KRAB-associated protein 1
	ATCTCTATCCC	. 1,00			alanyl-tRNA synthetase
	CACCGGACACT	1,00			cytochrome c oxidase subunit VIc
	GAATTTGTGTA	1,00		Hs.28707	signal sequence receptor, gamma
			-,		(translocon-associat
6554	AAATTAAAACA	1,00	0,45	Hs.198307	von Hippel-Lindau binding protein 1
6555	TTTTTCAATCA	1,00			KIAA0036 gene product
6556	CCTCTGGAGGC	1,00			P450 (cytochrome) oxidoreductase
6557	CCTCCCCTGCA	1,00			ESTs
6558	CATCCAAAACA	3,00			heterogeneous nuclear ribonucleoprotein
		, ,			H1 (H)
6559	GTCTCATTTGA	2,00	0,89	Hs.92381	nudix (nucleoside diphosphate linked
					moiety X)-type
6560	ACACCTCTAAA	2,00	0,89	Hs.273230	hypothetical protein FLJ10830
6561	ATGGAGACTTC	2,00	0,89	Hs.239760	citrate synthase
6562	TGATCACCTAT	2,00	0,89	Hs.177861	CGI-110 protein
6563	GGAATAAATTA	4,00			cytochrome c-1
6564	AACTGTCCTTC	3,00	1,30	Hs.194673	phosphoprotein enriched in astrocytes 15
6565	ATCTTGTTACT	9,00			fibronectin 1
	ATGGCTGGTAT	61,00	20,14	Hs.182426	ribosomal protein S2
6567	TTCTCTTCTCC	1,00	0,46	Hs.6106	RNA binding motif protein 4
IO COOL	TACACTGCTTT	1,00	0.46	Hs 284233	NICE-5 protein

6569	ATGGATGCACT	1,00	0.46	Hs.24983	hypothetical protein from EUROIMAGE
					2021883
	ACAAACAAAAG	1,00		Hs.23964	sin3-associated polypeptide, 18kD
6571	GCTCCAGCCAT	1,00	0,46	Hs.1706	interferon-stimulated transcription factor
					3, gamma
	ACCTCTGGCTT	1,00			paired mesoderm homeo box 1
6573	GGATGTAGAGA	1,00			huntingtin interacting protein 2
6574	CTTTCAGATGT	4,00	1,69	Hs.99910	phosphofructokinase, platelet
6575	GAGGGTGCCAA	2,00	0,91	Hs.8986	complement component 1, q
					subcomponent, beta polypep
	GCTTCCATCTT	3,00	1,32	Hs.55296	HLA-B associated transcript-1
	GCACCTCAGCC	3,00		Hs.10702	hypothetical protein DKFZp761H221
	CCTACAGATAA	1,00	0,47		heat shock 60kD protein 1 (chaperonin)
6579	AAGGAGCGGGA	1,00	0,47	Hs.43543	suppressor of white apricot homolog 2
6580	CCAAGGAATGG	1,00	0,47	Hs.271934	Homo sapiens mRNA; cDNA
					DKFZp434M162 (from clone DKF
6581	ATTCAGCCACG	1,00			KIAA1366 protein
6582	AGTTTTATTTG	1,00	0,47	Hs.236494	RAB10, member RAS oncogene family
6583	TGAGGAAGACA	1,00	0,47	Hs.159627	death associated protein 3
6584	AGGTCCCTGTC	1,00	0,47	Hs.109571	translocase of inner mitochondrial
					membrane 10 (yeas
	CGCTTTGCGCG	1,00			valosin-containing protein
6586	AAGAAGATAGA	16,00	5,89	Hs.184776	ribosomal protein L23a
6587	CTGAGACAAAG	8,00	3,16	Hs.101025	basic transcription factor 3
6588	GAAACAAGATG	6,00	2,45	Hs.78771	phosphoglycerate kinase 1
6589	GTGCGCTGAGC	6,00			major histocompatibility complex, class I,
6590	TGTGATCAGAC	13,00	4.90	Hs.107476	ATP synthase, H+ transporting,
	•	· 1			mitochondrial F1F0, s
6591	GTGATGGTGTA	6,00	2,46	Hs.197345	thyroid autoantigen 70kD (Ku antigen)
6592	TTTTGTGTGAA	3,00			hypothetical protein FLJ10024
6593	GAGTGAGTGAG	2,00	0,93	Hs.10463	ESTs, Weakly similar to C44C1.2 gene
		·			product [C.eleg
6594	GGAGGGATCAG	3,00	1,34	Hs.6196	integrin-linked kinase
6595	GTTAACTGGGA	1,00	0,47	Hs.48295	RNA helicase family
6596	GCAGGAACAGC	1,00	0,47	Hs.4770	KIAA1068 protein
6597	TAAGTTCCTTC	1,00	0,47	Hs.237971	ESTs
6598	TGTACATTCTG	1,00	0,47	Hs.1624	ephrin-A1
6599	CGGGATGCAGA	1,00	0,47	Hs.155560	calnexin
6600	AAGATTGGGGT	2,00	0,94	Hs.169610	CD44 antigen (homing function and
					Indian blood group
6601	GACTCACTTTT	13,00	4,99	Hs.699	peptidylprolyl isomerase B (cyclophilin B)
6602	GTGGTACAGGA	6,00	2,50	Hs.31731	peroxiredoxin 5
	CCAAGTTTTTT	4,00	1,75	Hs.75914	coated vesicle membrane protein
6604	CCTTGACCAAT	2,00		Hs.8148	selenoprotein T
	GAATCATTTTG	2,00		Hs.75249	ADP-ribosylation factor-like 6 interacting
			,		protein
6606	CGCTGTGGGGT	2,00	0,95	Hs.7486	protein expressed in thyroid
	CGGGGAGATGA	2,00			N-myc downstream-regulated gene 2

6608	TCCGCGAGAAG	5,00	2,15	Hs.1665	zinc finger protein homologous to Zfp-36 in mouse
6609	TACAATAATTT	2,00	0,95	Hs.2730	heterogeneous nuclear ribonucleoprotein L
6610	CTGACCCCCTT	2,00	0,95	Hs.26492	beta-1,3-glucuronyltransferase 3 (glucuronosyltransf
6611	CTTTCCCCTTT	2,00	0,95	Hs.183800	Ran GTPase activating protein 1
6612	GGGCCCTTCCT	2,00	0,95	Hs.168073	DKFZP727M231 protein
6613	AACCAGTTTGT	2,00	0,95	Hs.15591	COP9 subunit 6 (MOV34 homolog, 34 kD)
6614	GAACTCAGGCC	1,00	0,48	Hs.83634	host cell factor C1 (VP16-accessory protein)
6615	CTGAGGCCTGG	1,00	0,48	Hs.82109	syndecan 1
6616	TTAGTTACCTT	1,00	0,48		ras homolog gene family, member A
6617	GGTAGCCCACG	1,00	0,48		trinucleotide repeat containing 5
6618	ACAGCTTTGTA	1,00	0,48		f-box and WD-40 domain protein 1B
6619	TTCAGGGCTTC	1,00	0,48	Hs.184242	sterol-C5-desaturase (fungal ERG3, delta-5-desaturas
Ĺ	GGAATGTACGT	9,00	3,64	Hs.429	ATP synthase, H+ transporting, mitochondrial F0 comp
6621	TCTCCAGGAAC	4,00			CGI-69 protein
6622	AAGTGAGGAGA	4,00	1,77	Hs.231840	WW domain binding protein 2
6623	CTGGCGCCGAT	3,00	1,38	Hs.183180	hypothetical protein
6624	GTGTGGTGGTG	2,00			GDP dissociation inhibitor 2
6625	TGGAAGGGCTC	2,00		Hs.31334	putative mitochondrial outer membrane protein import
6626	TAGTAAGTCAT	1,00	0,49		ribosomal protein L22
6627	TCTCCCTTCAA	1,00	0,49		CDC20 (cell division cycle 20, S. cerevisiae, homolo
6628	GGCAGATAGGT	1,00	0,49		Homo sapiens cDNA FLJ20115 fis, clone COL05594
6629	CAGATTGTGAA	1,00	0,49	Hs.142653	ret finger protein
	GACCCTGGGGA	1,00			ESTs, Weakly similar to Y063_HUMAN HYPOTHETICAL PROT
	TGGCGGAGTCC	1,00			ESTs, Weakly similar to F-box protein FBL9 [H.sapien
	GATGGTCAGTC	1,00	0,49	Hs.108779	DKFZP586E1519 protein
_	GGCTTGGTTTA	1,00	0,49	Hs.107362	KIAA0909 protein
6634	TAGCTCTATGG	4,00		Hs.190703	ATPase, Na+/K+ transporting, alpha 1 polypeptide
	GAATTTTATAA	4,00	1,82	Hs.202	benzodiazapine receptor (peripheral)
	CGGCTGAATTC	3,00		Hs.75888	phosphogluconate dehydrogenase
	TGGGTGAGCCA	7,00	3,01	Hs.249982	cathepsin B
6638	GAAGGCATCCT	3,00		Hs.250758	proteasome (prosome, macropain) 26S subunit, ATPase,
	GGGCCCCGCAG	2,00	0,99		KIAA0123 protein
6640	GCCCGCAGGGT	2,00		Hs.74375	dishevelled 1 (homologous to Drosophila dsh)
6641	GACAGATGGAC	1,00	0,50	Hs.83575	KIAA1533 protein

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6642	GAGTAGAGGCC	1,00	0,50	Hs.77813	sphingomyelin phosphodiesterase 1, acid lysosomal (a
6643	GCAGCCCGCGG	1,00	0,50	Hs.71472	ESTs, Highly similar to unnamed protein product [H.s
6644	GTGATTGTTCA	1,00	0,50	Hs.6727	Ras-GTPase activating protein SH3 domain-binding pro
6645	CTCCTGAAGGC	1,00	0,50	Hs.4890	ubiquitin-conjugating enzyme E2E 3 (homologous to ye
6646	GATAGGTCGGG	1,00	0,50	Hs.154721	aconitase 1, soluble
6647	TTGTTTGTAAA	1,00	0,50	Hs.15020	homolog of mouse quaking QKI (KH domain RNA binding
6648	GTATAATTTGT	1,00	0,50	Hs.107979	small membrane protein 1
6649	AAAAATAAAGG	7,00	3,02	Hs.155101	ATP synthase, H+ transporting, mitochondrial F1 comp
6650	GCCTGTATGAG	26,00	10,09	Hs.180450	ribosomal protein S24
6651	TTTCTGTATGT	2,00			H3 histone, family 3B (H3.3B)
6652	TCTTGTGCATA	8,00	3,45	Hs.2795	lactate dehydrogenase A
6653	TCATCTTCAAC	2,00	1,00	Hs.75525	calreticulin
6654	TCACTTTCTTT	1,00	·	Hs.78865	TATA box binding protein (TBP)- associated factor, RN
	TGGAGGGGCCA	1,00		Hs.74362	ClpP (caseinolytic protease, ATP- dependent, proteoly
6656	ATTGTGAACAA	1,00			ESTs
6657	TCTGTTTCCAG	1,00		Hs.155227	
6658	TGGAATGAGCG	1,00	0,51	Hs.153998	creatine kinase, mitochondrial 1
					(ubiquitous)
	GCTGGAGCGCC	1,00		Hs.12284	ESTs, Weakly similar to HS6B_DROME HEAT SHOCK PROTEI
6660	GCTTTTTAGAA	7,00			high-mobility group (nonhistone chromosomal) protein
6661	AAGTTTCCAAT	2,00	1,01	Hs.2903	protein phosphatase 4 (formerly X), catalytic subuni
6662	TTTCCTTCCTT	2,00	1,01	Hs.104143	clathrin, light polypeptide (Lca)
	CCTTCCAAATT	4,00			malate dehydrogenase 2, NAD (mitochondrial)
	TCCTCTTTCCA	1,00			natural killer cell transcript 4
	TGTCCTGGTTG	1,00			lipoma HMGIC fusion partner
	GTGGAAGACGA	1,00		Hs.80395	mal, T-cell differentiation protein
6667	ATTTGCCTCTG	1,00	0,52	Hs.7393	Homo sapiens mRNA full length insert cDNA clone EURO
6668	CTGGGCCTCTG	1,00	0,52	Hs.50868	solute carrier family 22 (organic cation transporter
6669	TCTGAATCGGG	1,00	0,52	Hs.4316	trinucleotide repeat containing 12
6670	GTGTGAAATAA	1,00	0,52	Hs.199179	RAN binding protein 2
6671	CACGGACTCGT	1,00	0,52	Hs.170195	bone morphogenetic protein 7 (osteogenic protein 1)
	GCAGCAGGAAG	1,00	0,52	Hs.165743	tumor suppressing subtransferable candidate 4
6673	CGGGTAGTATT	1,00	0,52	Hs.1437	glucosidase, alpha; acid (Pompe disease, glycogen st

6674	AAAATATTTTA	1,00	0,52	Hs.119000	actinin, alpha 1
	TTTGTTAATTC	3,00			heterogeneous nuclear ribonucleoprotein
			•		H2 (H')
6676	GTGCTGAATGG	27,00	10.97	Hs.77385	myosin, light polypeptide 6, alkali,
		,	,		smooth muscle a
6677	GGCACCTCTGC	2,00	1.03	Hs.3763	amyloid beta (A4) precursor protein-
		,	.,		binding, family
6678	GGTGCTGGAGA	2,00	1,03	Hs.155020	putative methyltransferase
	AATGCTTTGTT	10,00			Tubulin, alpha, brain-specific
	CGGAGTCCATT	4,00			neural precursor cell expressed,
		]			developmentally dow
6681	CGATGGTCCCC	3,00	1.50	Hs.7771	B-cell associated protein
	CAGCTCCGCTT	2,00		Hs.82113	dUTP pyrophosphatase
	ATAAAAAAAA	1,00		Hs.83942	cathepsin K (pycnodysostosis)
	GGACCTGCGCC	1,00		Hs.8297	ribonuclease 6 precursor
	GAAACCCTCAC	1,00		Hs.75859	chromosome 11 open reading frame 4
	GCAACCACGAC	1,00			CGI-36 protein
	GGATGCATTAG	1,00		Hs.172635	
	GCCAGCGTCAT	1,00			gamma-tubulin complex protein 2
	CCAAACGTGTA	12,00			H3 histone, family 3A
	TTCACAAAGGA				
0090	TICACAAAGGA	2,00	1,05	HS. 70913	proteasome (prosome, macropain)
6601	GGTTTGGCTTA	7.00	2.25	Un 72010	subunit, alpha type,
0091	GGIIIGGCIIA	7,00	3,25	Hs.73818	ubiquinol-cytochrome c reductase hinge
6602	GGTTGGCAGGG	5,00	2.41	Un 2745	protein
	TATAATCTTTA			Hs.3745	milk fat globule-EGF factor 8 protein
		1,00		Hs.923	single-stranded DNA-binding protein
	TAAAATTGCTG CCGGAATGTGG	1,00		Hs.75257	Hairpin binding protein, histone
0095	CCGGAATGTGG	1,00	့ ပ,၁၁	Hs.54702	xylosylprotein beta1,4-
6606	TTCTATTCCAC	1.00	0.50	11- 2024	galactosyltransferase, polype
0090	TTGTATTCCAG	1,00	0,53	Hs.3631	immunoglobulin (CD79A) binding protein
6697	GTACGTCTGGC	1,00	0.53	Hs 279754	pilin-like transcription factor
	AAAACTTTGTC	1,00	0.53	Hs 272458	protein phosphatase 3 (formerly 2B),
		1,00	0,00		catalytic subun
6699	GGGAGTAATAG	1,00	0.53	Hs.26045	protein tyrosine phosphatase, receptor
		.,,,,,	2,00		type, A
6700	TTTAATTTGTA	1,00	0.53	Hs.182793	golgi membrane protein GP73
	AATACTTTTGT	1,00			DKFZP564M2423 protein
	CATTTCAGAGA	1,00			BCL2-associated athanogene 3
	TCAATAAAGGA	1,00			ubiquitin-conjugating enzyme E2D 3
"	. 5, 5 1, 7 5 10 0/1	.,55	5,55		(homologous to ye
6704	GGAACTTTTAG	3,00	1 53		similar to glucosamine-6-sulfatases
	GGGGCAGGGCC.	12,00			eukaryotic translation initiation factor 5A
	TTCTAACATAT	2,00			ATPase, Na+/K+ transporting, beta 1
اک کا		2,00	1,00	1 13.7 0028	polypeptide
6707	GGGAGGGAAGA	2,00	1.06	Hs.75243	
	AAGATCCCCGC	5,00			bromodomain-containing 2
	GTATTGGCCTT				divalent cation tolerant protein CUTA
		2,00			transmembrane 9 superfamily member 2
0/10	TTCTCCCGCTT	2,00	1,07	⊓S. 110126	protective protein for beta-galactosidase

	Γ				(galactosi
6711	GAGTAAAAAAT	1,00	0.54	Hs.944	glucose phosphate isomerase
	TACAATAAACC	1,00		Hs.9071	progesterone membrane binding protein
_	TTGACCCTGGG	1,00		Hs.8128	phosphatidylserine decarboxylase
	AGCCTGACTGC	1,00		Hs.80206	glucose-6-phosphate dehydrogenase
	GAGGATTTGGG	1,00		Hs.43549	uncharacterized hematopoietic
00	0,100,1111000	',00	0,0 .	1.0. 100 10	stem/progenitor cells
6716	TGCTTTCAAAA	1,00	0.54	Hs 31439	serine protease inhibitor, Kunitz type, 2
	AAAGCAGTTTA	1,00		Hs.262962	
	CTGTACTAGGT	1,00			nesca protèin
	TGGGAGAAGTG	1,00		Hs.184544	
	TCCTTGTTGGC	1,00			glyoxylate reductase/hydroxypyruvate
0,20	10011011000	1,00	0,04	110.100742	reductase
6721	GAGACTGCTTC	1,00	0.54	Hs.118078	
	ACAACTCAATA	4,00			brain protein I3
	CGCTTTTGTAG	2,00			DKFZP564A2416 protein
	GAGGTCCCTGG	5,00			proteasome (prosome, macropain)
0124	6766166166	3,00	2,40	115.74077	subunit, alpha type,
6725	ATTGTTTATGG	12,00	5 52	He 181163	high-mobility group (nonhistone
0/23	AIIGIIIAIGG	12,00	3,32	118.101103	chromosomal) protein
6726	CTGAGGCGCTT	1,00	0.55	Hs.78769	thimet oligopeptidase 1
	TTCATTTGTCT	1,00			solute carrier family 20 (phosphate
0121	TICATITOTO	1,00	0,55		transporter), me
6728	GAGAGCACCCT	1,00	0.55		postmeiotic segregation increased 2-like
0120	GAGAGCACCCI	1,00	0,55	115.273237	postificione segregation increased 2-like
6720	AGTGGCTGCCC	1,00	0.55	Hs.24435	ESTs
	CCCTTCACTGG	1,00		Hs.22557	ESTs, Highly similar to unnamed protein
0730	CCCTTCACTGG	1,00	0,55	П8.2255 <i>1</i>	product [H.s
6731	GCCTGGGACTC	3,00	1 58	Hs.98057	ESTs, Weakly similar to I68667
0/51	GOOTGOOACTC	3,00	1,50		transcription factor
6732	CACTCAGTGTG	2,00	1 10		solute carrier family 1 (glial high affinity
0102		2,00	1, 10	113.70079	glutama
6733	TTTCTAGGGGT	3,00	1 59	Hs 108969	PTD008 protein
	TCCAAGGAAGG	2,00			peroxisomal D3,D2-enoyl-CoA
0,04		2,00	1,10	113.10200	lisomerase
6735	GCAAAACCAGC	3,00	1.60	Hs.15071	chaperonin containing TCP1, subunit 8
0,00	00/100/100/100	0,00	1,00		(theta)
6736	GCTGTTTTGTT	1,00	0.56		Leman coiled-coil protein
	GAGCTGTTGGT	1,00			integrin, alpha E (antigen CD103, humar
3, 37	5,1001011001	1,00	0,50	1 13.00 1	mucosal lymp
6738	CATAGAGCCAC	1,00	0.56	Hs.6118	hypothetical protein FLJ10968
	AAACCTCTCAA	1,00			DKFZP586M1523 protein
	GAGGTGCTCTA	1,00			uncharacterized hematopoietic
J, 40	S. COTOCIOIA	,,00	0,50	1 13. 10 1303	stem/progenitor cells
6741	AGCCCGCCGCG	1,00	0.56	He 15/026	tumor suppressing subtransferable
0741		1,00	0,50	11 13. 134030	candidate 3
6742	CTAACTTCGTT	1,00	0.56	Hs.14838	hypothetical protein FLJ10773
	GGCTTTACCCT			He 110140	eukaryotic translation initiation factor 5A
	CATCTGTGAGC	9,00			
01 <del>44</del>	UNIUIGIGAGG	2,00	1,11	11 15.70 109	death-associated protein

6745	CCCCCAATTCT	2,00	1,11	Hs.194534	vesicle-associated membrane protein 2 (synaptobrevin
6746	AATTTCTATTT	2,00	1,12	Hs.5322	guanine nucleotide binding protein (G protein), gamm
6747	GCGATTCCGGA	2,00	1,12	Hs.283724	ESTs, Weakly similar to HYA22 [H.sapiens]
6748	TCAGTTTGGAG	3,00	·	Hs.3873	palmitoyl-protein thioesterase 1 (ceroid- lipofuscino
6749	TGTCAGAGATG	1,00	0,57	Hs.73957	RAB5A, member RAS oncogene family
6750	ATTGCTTTTGA	1,00	0,57	Hs.40500	similar to S. cerevisiae RER1
6751	GCTAGTGATGT	1,00	0,57	Hs.284162	60S ribosomal protein L30 isolog
6752	CCGGCGCGTGT	1,00	0,57	Hs.107387	CGI-20 protein
6753	TGCACCACAGA	2,00	1,13	Hs.9534	signal peptidase complex (18kD)
6754	ATGGCCAACTT	2,00	1,13	Hs.227835	KIAA1049 protein
6755	GTGTCTCATCT	2,00	1,13	Hs.144904	nuclear receptor co-repressor 1
_	ATGAGCTATGA	1,00			transmembrane protein 4
6757	GTCAGAACTTG	1,00		Hs.82101	pleckstrin homology-like domain, family A, member 1
6758	CTGAAATTCGG	1,00	0,57	Hs.79658	casein kinase 1, epsilon
6759	ACAAATTATGA	1,00	0,57	Hs.78902	voltage-dependent anion channel 2
6760	TTTGCAATTAT	1,00	0,57	Hs.75137	KIAA0193 gene product
6761	GGGCTGGGGTA	2,00	1,14	Hs.90436	sperm associated antigen 7
	GTGGGGCTAGG	2,00			protein phosphatase 5, catalytic subunit
	GTGTGTAAAAA	2,00			accessory proteins BAP31/BAP29
6764	AATGGATTACC	1,00		Hs.82202	ribosomal protein L17
$\overline{}$	CTTGTAACAGA	1,00			laminin, beta 1
	GAAGGCATCTT	1,00		Hs.71377	p138-tox
	GGGGCCCCGT	1,00		Hs.5809	putative transmembrane protein
-	TGATGCGCGCT	1,00			tumor suppressor deleted in oral cancer- related 1
6769	TCTGTGACCTT	1,00	0,58	Hs.198793	KIAA0750 gene product
6770	AGACAAGTTTA	1,00		Hs.120874	
6771	TTCCTGACTAC	1,00	0,58	Hs.10098	putative nucleolar RNA helicase
6772	GCAATAAATGG	2,00			drebrin 1
6773	AGGAGCGGGGT	2,00	1,16	Hs.252189	syndecan 4 (amphiglycan, ryudocan)
6774	TCAGAAGTTTT	2,00		Hs.243901	Homo sapiens cDNA FLJ20738 fis, clone HEP08257
6775	GCGGGGTACCC	4,00	2,18	Hs.227823	pM5 protein
6776	GCGGGAGGGCT	3,00			ADP-ribosylation factor-like 2
-	AAAGCAAACCA	2,00			hypothetical protein FLJ20159
	GCACCCTCAGA	1,00			regulator of G-protein signalling 3
	ATCTTTTAAAA	1,00		Hs.75867	solute carrier family 20 (phosphate transporter), me
6780	GGGCAGAATTG	1,00	0,59		KIAA0370 protein
	ACTITITAAAA	1,00			diaphanous (Drosophila, homolog) 2
	TATTCTCAATA	1,00			asparaginyl-tRNA synthetase
	CGCTGTGTGCT	1,00			glucosidase, beta; acid, pseudogene
	TCTGGACTCGG	1,00			ATP-binding cassette, sub-family F
L					(GCN20), member 2

PCT/EP01/15179

6705	TTTCCTACATC	1.00	0.50	110 407747	DVEZDECCOMA protein
	TTTCGTAGATG	1,00			DKFZP566C243 protein
	ACAAATCCTTG	3,00			FK506-binding protein 1A (12kD)
	GTCTTAACTCA	2,00		Hs.5074	similar to S. pombe dim1+
6788	GAGTTATGTTG	2,00	1,17	Hs.279915	translocase of inner mitochondrial
					membrane 8 (yeast
6789	GAGTAGAGAAA	3,00	1,71	Hs.145279	SET translocation (myeloid leukemia-
					associated)
	ATAGGTCAGAA	4,00			KIAA0911 protein
-	TAATGGTAACT	4,00			cytochrome c oxidase subunit Va
6792	GCAGTGGCCTC	2,00	1,19	Hs.184276	solute carrier family 9 (sodium/hydrogen
				٠.	exchanger),
	AACATTCTAAG	1,00			syntaxin binding protein 3
-	TGAGAGACATC	1,00			Mulibrey nanism
6795	TGCTGCCTGTT	3,00	1,74	Hs.118110	bone marrow stromal cell antigen 2
6796	TGGATCCTAGA	3,00	1,75	Hs.5273	NADH dehydrogenase (ubiquinone) Fe-S
					protein 3 (30kD
6797	GATCACAGTTT	3,00	1,75	Hs.234489	lactate dehydrogenase B
6798	AATGGAAATCG	1,00			hepatocellular carcinoma associated
	•				protein; breast
6799	TTAATATATGC	1,00	0,61	Hs.237955	hypothetical protein PRO2706
6800	TTTTAAACTTG	1,00	0,61	Hs.226770	DKFZP566C0424 protein
6801	CCCCAGTCGGC	1,00		Hs.171596	
6802	CTGGGTGAAGT	1,00		Hs.169388	
	ATCAAGAATCC	1,00			interferon, gamma-inducible protein 30
	TAATAAATGCT	1,00			(Manual assignment) MEMOREC GS2L1
		.,,,,	0,0 1		putative phospholi
6805	ATTATTTTTCT	7,00	3 79	Hs.153	ribosomal protein L7
	ATTCCAATCTT	2,00			clathrin, heavy polypeptide (Hc)
	ACTGGGTCTAT	10,00			non-metastatic cells 2, protein (NM23B)
		.0,00	0,0 .	110.270100	expressed in
6808	TTAAGAGGGGG	2,00	1 22	Hs 243886	nuclear autoantigenic sperm protein
		_,00	.,	10.2 1000	(histone-binding
6809	CAAAGACAATT	1,00	0.62	Hs 23528	HSPC038 protein
	ACCCTCTCCCT	1,00			eukaryotic translation initiation factor 4E-
00.0	/.0001010001	1,00	0,02	113.10122	like 3
6811	TGGACCAGGCG	1,00	0.62	Hs 167641	hypothetical protein from EUROIMAGE
0011	100/100/1000	1,00	0,02	113.107041	1703145
6812	ATGGTGCTGAC	1,00	0.62	Hs.166	sterol regulatory element binding
0012	7110010010710	1,00	0,02	113,100	transcription fact
6813	CTTTGATCAGG	1,00	0.62	He 118249	brefeldin A-inhibited guanine nucleotide-
00.0	0.110/110/100	1,00	0,02	110.1702-10	exchange pr
6814	GATGACGACTC	1,00	0.62	Hs 107019	symplekin; Huntingtin interacting protein I
	ACAAACTTAGG	6,00			calmodulin 1 (phosphorylase kinase,
		5,55	5,50	1 13. 17 7 000	delta)
6816	TCTTTACTTGA	2,00	1 22	Hs.6895	actin related protein 2/3 complex, subunit
33.13	JULIA	2,00	1,23	119.0080	3 (21 kD)
6817	AGTTGAAATTC	2,00	1 23	He 180429	KIAA1181 protein
	TCCGGCCGCGA	4,00			hypothetical protein
	ATGCGAAAGGC				<del></del>
0019	A I GCGAAAGGC	2,00	1,24	Hs.89466	dodecenoyl-Coenzyme A delta

					linewayana (2.2 trans. and
0000	444744047	0.00	4.04	11- 700	isomerase (3,2 trans-eno
	AAATAAAGAAT	2,00		Hs.790	microsomal glutathione S-transferase 1
6821	GTGCCTGTGCT	1,00	0,63	Hs.8888	Homo sapiens mRNA; cDNA
2000	T100100T00	4 00	0.00	11- 70070	DKFZp434C2019 (from clone DK
	TAGGACCCTGC	1,00		Hs.76873	hyaluronoglucosaminidase 2
	GTAATGAAGCA	1,00		Hs.5011	RNA binding motif protein 9
$\overline{}$	CCGAAAAAGTG	1,00		Hs.24763	RAN binding protein 1
	CTTCAGGACCT	1,00			syntaxin binding protein 1
	GACACAGGCAG	1,00		Hs.143641	low density lipoprotein receptor-related protein 3
6827	CAAATAAAAAG	1,00	0,63	Hs.1116	lymphotoxin beta receptor (TNFR superfamily, member
6828	GTGAAAAAAAA	2,00	1,25	Hs.181373	accessory proteins BAP31/BAP29
6829	GCAGCTCAGGC	3,00	1,82	Hs.79572	cathepsin D (lysosomal aspartyl
					protease)
6830	TTGACACTTTC	3,00			ESTs
6831	AACTAATACTA	7,00	3,97	Hs.118724	DR1-associated protein 1 (negative cofactor 2 alpha)
6832	GGTGATGAGGA	2,00	1,26	Hs.12107	putative breast adenocarcinoma marker (32kD)
6833	TAAGTTTAATT	1,00	0,64	Hs.75760	sterol carrier protein 2
	AGACCATATTA	1,00			sin3-associated polypeptide, 18kD
	TGGGCAGCTGG	1,00			ribosomal protein S9
-	ATATTTTCCTG	1,00			Novel human gene mapping to
		,,,,,,	-,-		chomosome 1
6837	ATGCCCGTGAG	1,00	0,64	Hs.118223	microfibrillar-associated protein 4
	AGCTCTTGGAG	3,00			selenium binding protein 1
	GGCCAAAGGCC	2,00			KIAA0064 gene product
	ATCTGAAGCAA	2,00			granin-like neuroendocrine peptide
			·		precursor
6841	TAGCATTTTAA	1,00	0,64	Hs.77665	KIAA0102 gene product
6842	ATGTGGCACAT	1,00		Hs.67776	
6843	AGTTTCTTGTC	1,00	0,64	Hs.246381	CD68 antigen
6844	AAGTATTGTGT	1,00	0,64	Hs.100688	Homo sapiens cDNA FLJ11279 fis, clone PLACE1009444,
6845	TTTTGGGGGCT	2,00	1,29	Hs.26683	ESTs, Weakly similar to lamin B receptor homolog TM7
6846	AGTGTCTGTGA	3,00	1,88	Hs.8867	cysteine-rich, angiogenic inducer, 61
	TCACCCACACC	26,00			ribosomal protein L23
	AAGATTGGTGG	5,00		Hs.1244	CD9 antigen (p24)
	TAGTTGAAGTC	6,00			ubiquinol-cytochrome c reductase
		-,	-,		binding protein
6850	ATACTTTAATC	2,00	1.29	Hs.79274	annexin A5
	ATGGCTGGGCT	1,00		Hs.752	FK506-binding protein 1A (12kD)
	TTCGCTTCCTG	1,00		Hs.6120	Homo sapiens mRNA; cDNA
		.,55	5,50		DKFZp434E146 (from clone DKF
6853	GAATGTAAGTA	1,00	0,65	Hs.6113	staufen (Drosophila, RNA-binding protein)
6854	сттгестетет	1,00	0.65	Hs 138617	thyroid hormone receptor interactor 12
2007	3.1.0010101	1,00	<del></del>	1	landing trouble terebrot litteractor 12

	3C3C3C4AC3C4CCC	1,00	0 65	Hs.11747	hypothetical protein EL 120201
	CCTGACGCTCC GGGCCTGACAT				hypothetical protein FLJ20391 Homo sapiens clone 24804 mRNA
0000	GGGCCTGACAT	1,00		Hs.11039	sequence
	GGAAGCACGGA	4,00			proteasome (prosome, macropain) 26S subunit, non-ATP
6858	TCCTTCTCCAC	3,00			actinin, alpha 1
6859	AAATAGATCCA	2,00			catenin (cadherin-associated protein), beta 1 (88kD)
	CAGCCCCTCTT	2,00		Hs.82503	H.sapiens mRNA for 3'UTR of unknown protein
	AATACCTCGTG	2,00		Hs.24220	hypothetical protein
6862	CTGGTCCTCCT	1,00	0,66	Hs.9739	ESTs
	ATCAAGTGGAC	1,00		Hs.79077	KIAA0233 gene product
6864	GCTTATGTTAA	1,00	0,66	Hs.75187	translocase of outer mitochondrial membrane 20 (yeas
6865	AGCTTATTGAG	1,00	0,66	Hs.273077	Human DNA sequence from clone RP3-417M14 on chromoso
6866	GGCTTCCTGGG	1,00	0,66	Hs.253041	EST
6867	GACATCGAGGC	1,00			ribosomal protein L10a
	TGATGTCTGGT	2,00			PMEPA1 protein
	GCTAAGGAGAT	5,00			ras-related C3 botulinum toxin substrate 1 (rho fami
6870	GCTGTTGCGCG	7,00	4,20	Hs.8102	ribosomal protein S20
	GCAGGGTGGGG	2,00			calmodulin 2 (phosphorylase kinase, delta)
6872	GGCTGCCCTGG	3,00	1,94	Hs.74566	dihydropyrimidinase-like 3
	CAGTGAACAAG	1,00		Hs.9825	CGI-128 protein
	GTGGCCCGCAG	1,00		Hs.96200	neighbor of A-kinase anchoring protein 95
6875	CTGCAGTTAGC	1,00	0,67	Hs.9614	ESTs, Weakly similar to Ps 2
	GAAAGGATTTT	1,00		Hs.84318	replication protein A1 (70kD)
	TGCATATCATC	1,00		Hs.79090	exportin 1 (CRM1, yeast, homolog)
6878	TGCGTCACCGT	1,00		Hs.78202	SWI/SNF related, matrix associated, actin dependent
	TCTTGATTTAC	1,00	0,67	Hs.74561	alpha-2-macroglobulin
	GTGATGGGGCC	1,00	0,67	Hs.62620	chromosome 6 open reading frame 1
6881	GGTGTGGGTGA	1,00		Hs.285363	ESTs
6882	GGCCACTCTAG	1,00			phenylalanine-tRNA synthetase-like
	CTCCCTTGCCC	1,00			CGI-120 protein
6884	GGCCCTGGTGT	1,00			cyclin G associated kinase
6885	AGGTTCTGCCT	1,00			putative nuclear protein
6886	TGATAATTCAA	5,00			Homo sapiens cDNA FLJ11279 fis, clone PLACE1009444,
6887	TTTTCTGCATA	1,00	0,68	Hs.77318	platelet-activating factor acetylhydrolase, isoform
	TGTTAGATTTC	1,00	0,68	Hs.77254	chromobox homolog 1 (Drosophila HP1 beta)
6889	TGGAAGAAACT	1,00	0,68	Hs.37810	ESTs
	GCCAGGTGGAA	1,00			Homo sapiens G protein beta subunit

					r =
					mRNA, partial cd
6891	TTTGTTTTG	1,00	0,68	Hs.183418	cell division cycle 2-like 1 (PITSLRE
					proteins)
6892	TTAAGAGGGAC	1,00	0,68	Hs.178137	transducer of ERBB2, 1
6893	TTGGCAACATT	1,00	0,68	Hs.11463	UMP-CMP kinase
6894	TCGAAGAACCG	2,00	1,36	Hs.76294	CD63 antigen (melanoma 1 antigen)
6895	CACTTTTGGGC	2,00			LIM and SH3 protein 1
	TTTCTGCACTT	1,00			ESTs, Weakly similar to ALU7_HUMAN
		1,55	,		ALU SUBFAMILY SQ
6897	CAGGGCGGGTT	1,00	0.69	Hs.23978	scaffold attachment factor B
	TAACCTGCTAT	1,00			cullin 1
	TACCATCAATA	41,00			glyceraldehyde-3-phosphate
	171007170701171	,00	20,00	110.100-110	dehydrogenase
6900	ACTGAGGTGCC	2,00	1 38	Hs 284159	FIBP-1 protein
	GCCATATTATG	1,00		Hs.19280	cysteine-rich repeat-containing protein
10301	OCCAIALIAIG.	1,00	0,70		S52 precurso
6002	ATTTTAGAATT	1,00	0.70		nucleosome assembly protein 1-like 1
	TTCAAAAAAAA				
		1,00			CGI-147 protein
169041	AGGAGCTGCTG	4,00	2,63		NADH dehydrogenase (ubiquinone) Fe-S
0005	00000704000				protein 8 (23kD
16905	GGGGGTCACCG	4,00	2,63		ATP synthase, H+ transporting,
					mitochondrial F0 comp
	CTGTTAGTGTG	4,00			malate dehydrogenase 1, NAD (soluble)
6907	CGCCGACGATG	11,00	6,80		interferon, alpha-inducible protein (clone
1 222					IFI-6-16)
	AAAAGGTTATG	1,00			sorting nexin 1
	TAAACCTGTCT	1,00			hypothetical protein
	GCTGTGCCTGG	1,00			protease, serine, 4 (trypsin 4, brain)
	TGCCCACTCAT	1,00			transmembrane 7 superfamily member 2
6912	CGGGATTCCTC	1,00	0,71		protective protein for beta-galactosidase
6012	TTOATOTAGAG	4.00	0.74		(galactosi
	TTGATGTACAG	1,00		Hs.11482	splicing factor, arginine/serine-rich 11
	TTTGGAAATCA	1,00			CGI-99 protein
6915	TTTTATTGGAA	1,00	0,71	Hs.106242	cytochrome P450, subfamily IVF,
0040			0.05		polypeptide 3 (leuko
6916	GGTAGCCTGGG	3,00	2,05	Hs.108327	damage-specific DNA binding protein 1
					(127kD)
	ATGCAGCCATA	2,00			ornithine decarboxylase 1
	AATTTTATTTC	2,00		Hs.2853	poly(rC)-binding protein 1
	CTTAAGGATTC	2,00			DKFZP564M2423 protein
6920	ACTGAGGAAAG	3,00	2,06	Hs.77326	insulin-like growth factor binding protein
					3
6921	TTTCTGAAAA	4,00	2,70	Hs.76136	thioredoxin
6922	GCTGCACCGGT	1,00	0,72	Hs.70582	ESTs, Moderately similar to AF144056_1
			·		apoptosis rel
6923	TACACGTGAGG	2,00	1,42	Hs.11156	hypothetical protein
	CTGGGCCTGGC	4,00		Hs.74573	similar to vaccinia virus HindIII K4L ORF
<b></b>	TTTGTAGATGG	2,00		Hs.3069	heat shock 70kD protein 9B (mortalin-2)
	GGGGGTTGGTT	1,00		Hs.4779	KIAA1150 protein
الاعتدا	22000110011	1,00	7,72	13.7113	INVALIDO PIOLEIR

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6927	TCCTGCAGCTG	1,00	N 72	Hs 283111	HCDI protein
	ATGTTAGGGAT	1,00	0,72	He 160206	vesicle-associated soluble NSF
0320	l	1,00	0,72	115.109200	attachment protein re
6020	GGGTCTGCGGG	1,00	0.72	Hs.146219	
	AACTTGATACG	1,00			stearoyl-CoA desaturase (delta-9-
0930	ACTIGATAGG	1,00	0,72	115.119391	desaturase)
6931	TGCATTAACTG	1,00	0,73	Hs.7351	cyclic AMP phosphoprotein, 19 kD
6932	TTAAACTCCAA	1,00		Hs.273369	uncharacterized hematopoietic
		,	·		stem/progenitor cells
6933	ACCTGCCGACA	1,00	0,73	Hs.25664	tumor suppressor deleted in oral cancer- related 1
6934	AAGGAGCTGGC	1,00	0.73	Hs.155165	zinc finger protein-like 1
6935	CAGGGTCCCCA	1,00			hypothetical protein FLJ10647
-	TCCGAGCCCCC	1,00			thioredoxin reductase beta
	TTAGATAAGCA	2,00		Hs.82916	chaperonin containing TCP1, subunit 6A (zeta 1)
6938	GTTACAAACTA	1,00	0.74	Hs.75248	topoisomerase (DNA) II beta (180kD)
	TTGGGAGTGAG	1,00			imidazoline receptor candidate
	CTGGTGATGGC	1,00		Hs.242463	
	TACGTACTGCC	1,00			translocase of inner mitochondrial
		, , ,	-,-		membrane 13 (yeas
6942	GATGCCTCTGC	1,00	0,74		ESTs, Weakly similar to TBP-associated
					factor [H.sap
	CTTCCAGCTAA	11,00			annexin A2
6944	TTGTTGTTGAA	6,00	4,14	Hs.182278	calmodulin 2 (phosphorylase kinase, delta)
6945	ACTGCTTGCCC	3,00	2,18		methionine adenosyltransferase II, alpha
6946	GCTGGGGTGGG	1,00	0,75	Hs.86131	Fas (TNFRSF6)-associated via death domain
6947	GGCAGGCACAA	1,00	0,75	Hs.75105	emopamil-binding protein (stérol
					isomerase)
_	CAGCCAAATAA	1,00			F-box protein FBG2
	AATAGCTCAGG	1,00		Hs.4990	KIAA1089 protein
	AAGAAGCAAGA	1,00		Hs.2953	ribosomal protein S15a
6951	TGTTGTGCGCG	1,00	0,75	Hs.11364	potassium inwardly-rectifying
6050	COTOTOOTTTO	1 00	0.70	11- 5000	channel,subfamily J, m
	CCTGTCCTTTC	1,00		Hs.5300	bladder cancer associated protein
6953	CCTATGTAAGG	1,00	0,76	HS.146381	RNA binding motif protein, X chromosome
6954	CTGGGCGTGTC	5,00	3,58	Hs.161554	hypothetical protein FLJ20159
6955	TTTCCAATCTC	1,00			vascular endothelial growth factor
6956	CTACCAGCACC	1,00			hypothetical protein FLJ20657
	GGGCTGGGCCC	2,00			6-phosphogluconolactonase
	AAAAAGCAGAT	3,00		Hs.75428	superoxide dismutase 1, soluble
					(amyotrophic lateral
	TTTTACAGTAC	1,00	0,78	Hs.86347	hypothetical protein
6960	CTGTGTAAGCT	1,00		Hs.79137	protein-L-isoaspartate (D-aspartate) O-methyltransfe
6961	CGGCGCTCCCT	1,00	0.78	Hs.29285	ZYG homolog
3331	2200010001	.,,00	3,,0		- i o nomony

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	GCATCTGTTTA	1,00			hypothetical protein dJ483K16.1
	CTCTGCTCGGC	1,00		Hs.241531	
	TAGGTTCGTGT	1,00			cysteinyl-tRNA synthetase
	TGAAGTAACAA	3,00			putative translation initiation factor
	GGGGAAATCGC	24,00			thymosin, beta 10
	TTGCTAGAGGG	2,00			ubiquitously-expressed transcript
	CAGACGCTCCG	1,00			CGI-139 protein
	TGGTGCAGCAT	1,00			30S ribosomal protein S7 homolog
6970	TCTGCAAATTA	1,00		l	Homo sapiens cDNA FLJ10433 fis, clone NT2RP1000478,
6971	TGAATGGCCTA	1,00			host cell factor homolog
$\overline{}$	GAAATGATGAG	5,00			prefoldin 5
	CGCCGCGGTGG	8,00		Hs.4835	eukaryotic translation initiation factor 3, subunit
	TTGTAAAAGGA	2,00			valosin-containing protein
6975	AGCAGGGCTCC	5,00	3,73	Hs.100623	phospholipase C, beta 3, neighbor
					pseudogene
	TGAGCCTCGTG	2,00			enolase 1, (alpha)
	GAAATACAGTG	1,00			5'(3')-deoxyribonucleotidase
	GATAATTTTTG	1,00			HSPC142 protein
	GGCGCCAAAAA	1,00			kinesin-like 4
	CCAATAAAGTG	1,00			retinol-binding protein 1, cellular
6981	ATCTTTCTGGC	4,00	3,05		tyrosine 3-monooxygenase/tryptophan 5-monooxygenase
	GCTGCGGTCCT	1,00		Hs.795	H2A histone family, member O
6983	CAATTACCTGC	1,00		Hs.284263	Homo sapiens clone H63 unknown mRNA
6984	CACGAAGATGC	1,00	0,81	Hs.10247	activated leucocyte cell adhesion molecule
6985	AGAACAAAACC	6,00	4,53		peroxiredoxin 1
	GGTGCCCAGTT	2,00	1,62	Hs.75607	myristoylated alanine-rich protein kinase C substrat
	GAGAGTGTCTG	3,00			tissue inhibitor of metalloproteinase 1 (erythroid p
:	GCACAAGAAGA	9,00		_ :	ATP synthase, H+ transporting, mitochondrial F0 comp
	TTTTATCTGGT	1,00		Hs.64239	Human DNA sequence from clone RP5-1174N9 on chromoso
$\overline{}$	TGTTTGGGGGC	1,00			ESTs
	ACTGTCTCCAG	1,00		Hs.16611	tumor protein D52-like 1
	ATGGCCAGAAA	1,00	0,82	Hs.141011	calmodulin 3 (phosphorylase kinase, delta)
	TATTTTTGTTA	1,00	0,82		thioredoxin reductase 1
	GGGGAGCTCGG	1,00			CGI-43 protein
6995	ACTGCCCCAAC	1,00		Hs.103147	ESTs, Weakly similar to similar to SP:YR40_BACSU [C.
6996	TGAATCTGGGT	2,00	1,63		SET translocation (myeloid leukemia- associated)
6997	ACCTCTCTAAA	1,00	0.83	Hs 94392	LDL induced EC protein

1 1

6998	GAGAGCTACAT	1,00	0,83	Hs.169919	electron-transfer-flavoprotein, alpha polypeptide (g
6999	AGGTAAAAGAG	1,00	0.83	Hs.103042	microtubule-associated protein 1B
_	GTGCTGCGTGA	2,00		Hs.4209	ribosomal protein, mitochondrial, L2
	GGGATGGCAGC	2,00			valyl-tRNA synthetase 2
	CCAAAAAAAA	2,00			hypothetical protein FLJ20455
	GCCCCAGGTAG	1,00		Hs.78466	proteasome (prosome, macropain) 26S
			<u> </u>		subunit, non-ATP
/004	AAACCTGGGAA	1,00	0,84	Hs.63788	propionyl Coenzyme A carboxylase, beta polypeptide
7005	GGGGGAGGGAA	1,00	0,84	Hs.108530	hypothetical protein FLJ10856
7006	TGTAAGTCTGC	2,00	1,66	Hs.119537	GAP-associated tyrosine phosphoprotein p62 (Sam68)
7007	ACGTGGTGATG	2,00	1,67	Hs.279945	HSPC023 protein
	CAGGACAGTTT	1,00			RAB2, member RAS oncogene family
-	ACCGTATTCCA	1,00			pyruvate kinase, muscle
	GCCGATCCTCG	2,00			tubulin-specific chaperone a
	TTGTCTGCCTT	3,00			Homo sapiens BAC clone CTB-122E10
		<i>'</i>	•		from 7q11.23-q21.1
7012	GAGAGTGTACT	1,00	0,85		translocase of inner mitochondrial
			,		membrane 17 (yeas
7013	GGGGGAATTTT	1,00	0,85		heterogeneous nuclear ribonucleoprotein K
7014	ATTTTTGGTGG	1,00	0.05	Hs.110	putative L-type neutral amino acid
1014	Ailinggigg	1,00	0,00		transporter
7015	ATCCGGCGCCA	4,00	3 26		transcription elongation factor B (SIII),
"		7,00	0,20	110.172772	polypeptid
7016	TCTGTCAAGAC	3,00	2 49	Hs.76572	ATP synthase, H+ transporting,
		0,00	Δ, ισ	110.7 007 12	mitochondrial F1 comp
7017	CGGACTCACTG	3,00	2,51	Hs.84700	similar to phosphatidylcholine transfer
					protein 2
7018	AAACCCCAATA	2,00	1,72	Hs.285501	Human rearranged immunoglobulin
					lambda light chain m
7019	TCAACTTCTGG	1,00	0,86	Hs.7594	solute carrier family 2 (facilitated glucose transpo
7020	GACCAGCCCAT	1,00	0,86	Hs.75799	protease, serine, 8 (prostasin)
	AGCAAGCCCCC	1,00			DAZ associated protein 1
7022	TTGTCGATGGG	1,00			hypothetical protein FLJ20442
	TACAGCACGGA	1,00			microsomal glutathione S-transferase 3
_	GAAACCGAGGG	4,00			hypothetical protein
	GTGTTAACCAG	15,00			ribosomal protein L15
	GTTGGTCTGTA	2,00			actin binding protein; macrophin
	· .		•		(microfilament and
7027	TCTTTCCCCAA	1,00	0,87	Hs.268557	pleckstrin homology-like domain, family
			•		A, member 3
7028	GTCTTTCTTGG	3,00	2,57	Hs.151536	RAB13, member RAS oncogene family
	CTCAGACAGTG	3,00			40S ribosomal protein S27 isoform
	GAACACATCCA	16,00			ribosomal protein L19
	GGGTTTTTATT	4,00		Hs.74497	nuclease sensitive element binding

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7000	ACAACCTTTCC	100	0.00	11-74040	protein 1
	AGAACCTTTGC	1,00			cytochrome c oxidase subunit VIc
	GCCTGATTTTT	1,00			HLA class II region expressed gene KE4
	GAGAGGTTGAT	1,00			paraneoplastic antigen MA1
	TGTAGTTTGAG	4,00			transcription elongation factor B (SIII), polypeptid
	AAGGAGATGGG	23,00	17,88	Hs.184014	ribosomal protein L31
	AACGCTGCCTG	2,00			adenine phosphoribosyltransferase
7038	CAGCCTCCCTG	1,00		Hs.75593	uroporphyrinogen III synthase (congenital erythropoi
7039	TTAATAGTGGG	1,00	0,89	Hs.18271	Homo sapiens mRNA; cDNA DKFZp434P1217 (from clone DK
7040	TGGCCCCACCC	12,00	9,68	Hs.198281	pyruvate kinase, muscle
7041	TGAAATAAAAC	9,00			nucleophosmin (nucleolar
					phosphoprotein B23, numatri
7042	TCCCTGGCTGT	1,00	0,90	Hs.78575	prosaposin (variant Gaucher disease and variant meta
7043	TATACCAATCA	1,00	0,90	Hs.75866	dimethylarginine
		·			dimethylaminohydrolase 1
7044	CAGATAACATA	1,00	0,90	Hs.75187	translocase of outer mitochondrial
					membrane 20 (yeas
	TGGATCCTCGT	1,00			hypothetical protein FLJ10479
	CCTATCAGTAA	1,00			microseminoprotein, beta-
7047	AGGTCAGAGGG	1,00			eukaryotic translation elongation factor 1 alpha 1
7048	CCTGCCCCTTC	1,00	0,90	Hs.173255	small nuclear ribonucleoprotein polypeptide A
7049	GGTACTCGATG	1,00	0,90	Hs.121576	aspartate beta-hydroxylase
7050	TTTGGAATGTT	2,00	1,80		matrin 3
	CTGGATCTGGG	2,00	1,80	Hs.75658	phosphorylase, glycogen; brain
7052	GTGCCATATTT	2,00		Hs.5337	isocitrate dehydrogenase 2 (NADP+),
7053	TACATCCGAAT	2,00	1,80	Hs.21321	Homo sapiens mRNA; cDNA DKFZp564E1363 (from clone DK
7054	GAAGTTATGAA	3,00	2,66	Hs.4112	t-complex 1
7055	TTTATTTCTAC	1,00	0,91	Hs.85963	CD36 antigen (collagen type I receptor, thrombospond
7056	TCAACGGTGTG	1,00	0,91		novel centrosomal protein RanBPM
7057	CCTCTTCAGGC	1,00			nucleophosmin (nucleolar phosphoprotein B23, numatri
7058	TCAGTTCTTGA	1,00	0.91	Hs.103804	heterogeneous nuclear ribonucleoprotein
		.,,,,,	5,51		U (scaffold
7059	CCTTTCCTTTA	3,00	2.67	Hs.74576	GDP dissociation inhibitor 1
	ACCGCCGTGGT	4,00		Hs.68877	cytochrome b-245, alpha polypeptide
-	GTAAAAGTTCT	1,00		Hs.75337	nucleolar phosphoprotein p130
	GTTTCAGGTAA	1,00		Hs.1526	ATPase, Ca++ transporting, cardiac muscle, slow twit
7063	TTCCAGACCTT	3,00	2.71	Hs.75722	ribophorin II
	TTATGGGATCT	13,00		Hs.5662	guanine nucleotide binding protein (G
ىت		, 0,00	, 0,00		Basimo hadicollac billaling protein (G

					In-tain) hate
7005	CTCCCCCCACC	4.00	0.00	11- 205720	protein), beta
	GTGGGGGGAGG	1,00			HLA class II region expressed gene KE2
	TAATTTTGGAT	1,00			CGI-43 protein
1067	GGGCAGATGCC	1,00	0,93	Hs.105894	Homo sapiens mRNA; cDNA
					DKFZp434G231 (from clone DKF
	GTTCTCCCACT	3,00			sec61 homolog
	AAGCCAGCCCC	3,00		Hs.1432	protein kinase C substrate 80K-H
	GGCCCCATTTT	1,00		Hs.88778	carbonyl reductase 1
	TAATATTTTTA	1,00			actinin, alpha 4
7072	GCGGGTGTGGA	1,00	0,95	Hs.78935	methionine aminopeptidase; eIF-2-
					associated p67
	CCCAGATGATA	1,00		Hs.7854	zinc/iron regulated transporter-like
	GGTTTGATTAC	1,00		Hs.69559	KIAA1096 protein
7075	TAATTACTCTT	1,00		Hs.44163	13kDa differentiation-associated protein
7076	TGTACTACTTA	1,00	0,95	Hs.286114	Homo sapiens cDNA FLJ10301 fis, clone NT2RM2000032
7077	AGGATGACCCC	1,00	0,95	Hs.25334	HSPC113 protein
7078	CTCTGTGTGGA	1,00			microtubule-associated protein, RP/EB
			•		family, member
7079	CTCTTATCACC	1,00	0.95	Hs.100043	ESTs, Weakly similar to PSF_HUMAN
		.,	-,		PTB-ASSOCIATED SPL
7080	ATAGACATAAA	2,00	1.90	Hs.78614	complement component 1, q
		_,			subcomponent binding prote
7081	TTTTCTGCTGG	2,00	1.90	Hs.204041	chromosome 14 open reading frame 3
	GTCTCACGTGG	1,00			Homo sapiens cDNA FLJ20187 fis, clone
		.,,,,	-,		COLF0433
7083	CACACAATGTG	1,00	0.96	Hs.117582	CGI-43 protein
	GATTGGGGATT	1,00	0.96	Hs. 103834	ESTs, Weakly similar to unnamed
	,	.,			protein product [H.s
7085	GCCCCTCCGGC	5,00	4,56	Hs.180859	16.7Kd protein
	AAAAAACTCCA	1,00			esterase D/formylglutathione hydrolase
	CTCTTCAGGAG	1,00		Hs.30954	phosphomevalonate kinase
	CAGAAGAGGCT	1,00			DiGeorge syndrome critical region gene
	,	.,	-,		6
7089	GGAACGGATGT	2,00	1.93	Hs.58589	glycogenin 2
-	GAGTGGGGGCT	2,00		Hs.14089	quiescent cell proline dipeptidase
	GAATCGGTTAT	3,00		Hs.80595	NADH dehydrogenase (ubiquinone) Fe-S
		-,			protein 5 (15kD
7092	AATATGCTTTA	1,00	0.98	Hs.77805	ATPase, H+ transporting, lysosomal
		,,,,,	-,		(vacuolar proton
7093	GTGTCTCGCAG	1,00	0.98	Hs.75510	annexin A11
	CTTAAATATCA	1,00			pyrophosphatase (inorganic)
	GTGGAGCGGAG	1,00			KIAA0890 protein
	ACAAACTGTGG	3,00		Hs.90370	actin related protein 2/3 complex, subunit
		5,55	_,00		1A (41 kD
7097	TATTTTGTGAG	2,00	1 97	Hs.75607	myristoylated alanine-rich protein kinase
		_,00	.,07		C substrat
7098	TGCATCTGGTG	5,00	4.72	Hs.75410	heat shock 70kD protein 5 (glucose-
		5,55	.,		regulated protein
	<u> </u>				In a Baracoa brocont

7099	GTGGCTGCTGT	1,00	1.00	Hs.41182	ESTs
	GAAAGAGCTGA	1,00			H2A histone family, member X
	GAGTCAGCATT	1,00			heterogeneous nuclear ribonucleoprotein
		.,,,,	.,		U (scaffold
7102	ATTGGCTTAAA	2,00	2,00	Hs.75323	prohibitin
	TTTGTTTTTGA	1,00			ESTs, Moderately similar to unnamed
	•		•		protein product
7104	ACCATTGGATT	1,00	1,01	Hs.146360	interferon induced transmembrane
					protein 1 (9-27)
7105	AGGCGAGATCA	4,00	3,87	Hs.233952	proteasome (prosome, macropain)
					subunit, alpha type,
7106	GGATTGTCTGG	2,00	2,01	Hs.83753	small nuclear ribonucleoprotein
					polypeptides B and B
	GGCGCCTCCTT	2,00			transaldolase 1
_	AAAATGTACTG	1,00			RAN binding protein 1
7109	TACTGTGGATC	1,00	1,02	Hs.21537	protein phosphatase 1, catalytic subunit,
					beta isofo
7110	ATGGTTAAAGG	1,00	1,02	Hs.142613	Homo sapiens cDNA FLJ10281 fis, clone
7444	0010000000	- 1 00	4.00	11 400040	HEMBB1001289
[/111]	CCAGCGTGGAA	1,00	1,02		ESTs, Highly similar to KINN_HUMAN
7440	TTOOAGGTGGG		4.05		NEURONAL KINESIN
[7112]	TTGGACCTGGG	5,00	4,85	Hs.89761	ATP synthase, H+ transporting,
7112	TTGGACTGAGC	2,00	2.04	Hs.6518	mitochondrial F1 comp ganglioside expression factor 2
	TGCCTTACTTT	1,00		Hs.80019	programmed cell death 6
	TTAGCAGTTGG	1,00			5'-3' exoribonuclease 2
	TGCAGGCCTGG	2,00			tryptophanyl-tRNA synthetase
	CCCCCTCGTGC	1,00		Hs.83636	adrenergic, beta, receptor kinase 1
	TAGTCTTAACA	1,00		Hs.75137	KIAA0193 gene product
	GCCCACATCCG	2,00		Hs.84753	KIAA0246 protein
	TGAGAGGGTGT	2,00			tyrosine 3-monooxygenase/tryptophan 5-
'		2,00	2,01		monooxygenase
7121	CCTTGGTGCCG	1,00	1.05		TNF receptor-associated factor 4
	ATTGTGAGGCC	1,00			sialyltransferase
	GAGAATCTGCT	1,00			cyclin B1
7124	CCGGGCGTGGT	1,00	1,05		delta-5 desaturase; delta-5 fatty acid
		, ,			desaturase
7125	ACTCCCTCCTT	1,00	1,06	Hs.76230	ribosomal protein S10
7126	TATCGTTGCCT	1,00	1,06	Hs.3887	proteasome (prosome, macropain) 26S
					subunit, non-ATP
	TGCTACTGGTA	1,00		Hs.3196	surfeit 1
	ATTCTTCGGAC	1,00			DKFZP434J154 protein
	TACCAGCACAG	1,00			nucleolar protein NOP5/NOP58
	AACTACATAGG	1,00		Hs.9527	apoptosis related protein APR-3
7131	TAACTGGAGGA	1,00	1,07	Hs.82906	CDC20 (cell division cycle 20, S.
					cerevisiae, homolo
7132	CCTGAACTGGC	1,00	1,07	Hs.74441	chromodomain helicase DNA binding
			<del></del>		protein 4
7133	CCGTAGTGCCT	1,00	1,07	Hs.6891	splicing factor, arginine/serine-rich 6

1,00	7134	AACTGCGGCAG	1,00	1,07	Hs.63908	Homo sapiens HSPC316 mRNA, partial
Cerevisiae -lik   Cerevisiae -lik   Geyruvoyl-tetrahydropterin   Synthase dimerization co   Synthase dimerization dimerial carrier, ade   Zerapiar in Chine   Certarior carrier, ade   Zerapiar in Chine   Certa						cds
Synthase/dimerization co	7135	TGTTCTCCATT	1,00	1,07	Hs.182255	cerevisiae)-lik
1,00	7136	TGTTTTATGT	1,00	1,08	Hs.3192	6-pyruvoyl-tetrahydropterin
						synthase/dimerization co
1,00	7137	CCTCCTCTGAC	1,00	1,08	Hs.28505	ubiquitin-conjugating enzyme E2H
7139   CCGGCCCTCC   1,00   1,08   Hs.155524   peanut (Drosophila)-like 2   7140   CAAGTTAGTGG   1,00   1,09   Hs.117582   CGI-43 protein   7141   GGGGGCGCTT   1,00   1,09   Hs.164280   solute carrier family 25 (mitochondrial carrier; ade   1,00   1,10   Hs.84981   X-ray repair complementing defective repair in Chine   7143   CTGCCTTCTTG   1,00   1,10   Hs.79081   protein phosphatase 1, catalytic subunit, gamma isof   7144   CGTACTGAGCG   1,00   1,11   Hs.180610   splicing factor proline/glutamine rich (polypyrimidi   1,11   Hs.6445   Homo sapiens (clone s153) mRNA fragment   1,11   Hs.2699   glycian 1   1,11   Hs.196176   enoyl Coenzyme A hydratase 1, peròxisomal   1,11   Hs.196176   enoyl Coenzyme A hydratase 1, peròxisomal   1,11   Hs.196176   enoyl Coenzyme A hydratase 1, peròxisomal   1,11   Hs.6551   AGCTGTGTAAA   1,00   1,12   Hs.6551   ATPase, H+ transporting, lysosomal (vacuolar proton   1,12   Hs.279923   hs.173714   MGRF-related gene X   7153   ATAGAGGCAAT   2,00   2,23   Hs.173714   MGRF-related gene X   7154   TGGTGACAGTT   2,00   2,23   Hs.285634   Homo sapiens HSPC222 mRNA, complete cds   1,00   1,13   Hs.33026   Human DNA sequence from clone RP3-402G11 on chromoso   1,14   Hs.285163   ESTs   1,00   1,14   Hs.285163   ES	L					
1,00						
1,09						
Carrier; ade						
7142 ATATAGGTCGT 1,00 1,10 Hs.84981 X-ray repair complementing defective repair in Chine protein phosphatase 1, catalytic subunit, gamma isof 7144 CGTACTGAGCG 1,00 1,10 Hs.180610 splicing factor proline/glutamine rich (polypyrimid) Homo sapiens (clone s153) mRNA fragment 7145 GACCCTGTCA 1,00 1,11 Hs.6445 Homo sapiens (clone s153) mRNA fragment 7146 GCTTGTTCTCT 1,00 1,11 Hs.2699 glypican 1 7147 CTGCAGACCCA 1,00 1,11 Hs.196176 enoyl Coenzyme A hydratase 1, peròxisomal 7148 CTGTGCATTTT 2,00 2,21 Hs.172207 non-POU-domain-containing, octamer-binding 100 1,12 Hs.279923 sulfotransferase family, cytosolic, 1C, member 2 ATPase, H+ transporting, lysosomal (vacuolar proton 7151 AGCTGTGAAA 1,00 1,12 Hs.279923 putative nucleotide binding protein, estradiol-induc 7152 CAAGATAAATT 1,00 1,12 Hs.177556 melanoma antigen, family D, 1 7153 ATAGAGGCAAT 2,00 2,22 Hs.173714 MORF-related gene X 7154 TGGTGACAGTT 2,00 2,23 Hs.9242 purine-rich element binding protein B 7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso dolichyl-phosphate mannosyltransferase polypeptide 3 hepatitis delta antigen-interacting protein 7158 ACGCAGGCCC 1,00 1,14 Hs.28163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.286761 licelf lip and palate associated transmembrane protei neurotrophic tyrosine kinase, receptor, 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor, 7163 TGAAACTCATC 2,00 2,27 Hs.85844	7141	GGGGGCGCCTT	1,00	1,09	Hs.164280	
repair in Chine						
gamma isof   gamma isof   gamma isof   splicing factor proline/glutamine rich   (polypyrimidi   factor proline sapparent   factor proline s	7142	ATATAGGTCGT	1,00	1,10	Hs.84981	
1,10	7143	CTGCCTTCTTG	1,00	1,10	Hs.79081	
7146 GCTTGTTCT 1,00 1,11 Hs.6445 Homo sapiens (clone s153) mRNA fragment 7146 GCTTGTTCT 1,00 1,11 Hs.2699 glypican 1 7147 CTGCAGACCCA 1,00 1,11 Hs.196176 encyl Coenzyme A hydratase 1, peroxisomal 7148 CTGTGCATTTT 2,00 2,21 Hs.172207 non-POU-domain-containing, octamer-binding 7149 AGTTTCCCAAT 3,00 3,27 Hs.75854 sulfotransferase family, cytosolic, 1C, member 2 7150 GGGTGCTTGGT 1,00 1,12 Hs.6551 ATPase, H+ transporting, lysosomal (vacuolar proton 7151 AGCTGTGTAAA 1,00 1,12 Hs.279923 putative nucleotide binding protein, estradiol-induc 7152 CAAGATAAATT 1,00 1,12 Hs.177556 melanoma antigen, family D, 1 7153 ATAGAGGCAAT 2,00 2,22 Hs.173714 MORF-related gene X 7154 TGGTGACAGTT 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds 7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,13 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.285163 ESTs 7161 TGGCCCCGGCC 1,00 1,14 Hs.285163 ESTs 1,00 1,14 Hs.285163 ESTs 1,00 1,14 Hs.285163 ESTs 1,00 1,14 Hs.285844 neurotrophic tyrosine kinase, receptor, 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor, 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7144	CGTACTGAGCG	1,00	1,10	Hs.180610	splicing factor proline/glutamine rich
	7145	GACCCCTGTCA	1.00	1.11	Hs.6445	
1,10			. ', ' '	.,		
7147 CTGCAGACCCA 1,00 1,11 Hs.196176 enoyl Coenzyme A hydratase 1, peròxisomal non-POU-domain-containing, octamerbinding sulfotransferase family, cytosolic, 1C, member 2 7149 AGTTTCCCAAT 3,00 3,27 Hs.75854 sulfotransferase family, cytosolic, 1C, member 2 7150 GGGTGCTTGGT 1,00 1,12 Hs.6551 ATPase, H+ transporting, lysosomal (vacuolar proton putative nucleotide binding protein, estradiol-induc melanoma antigen, family D, 1 7151 AGCTGTGAAA 1,00 1,12 Hs.177556 melanoma antigen, family D, 1 7153 ATAGAGGCAAT 2,00 2,23 Hs.9242 purine-rich element binding protein B 7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds 7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.285163 ESTs 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7146	GCTTGTTCTCT	1,00	1,11	Hs.2699	
peroxisomal   peroxisomal   peroxisomal     peroxisomal						
Dinding   Dinding   Sulfotransferase family, cytosolic, 1C, member 2   Member 2   ATPase, H+ transporting, lysosomal (vacuolar proton   Vacuolar proton				- 1	-	
AGTTTCCCAAT   3,00   3,27   Hs.75854   sulfotransferase family, cytosolic, 1C, member 2     T150   GGGTGCTTGGT   1,00   1,12   Hs.6551   ATPase, H+ transporting, lysosomal (vacuolar proton     T151   AGCTGTGTAAA   1,00   1,12   Hs.279923   putative nucleotide binding protein, estradiol-induc     T152   CAAGATAAATT   1,00   1,12   Hs.177556   melanoma antigen, family D, 1     T153   ATAGAGGCAAT   2,00   2,22   Hs.173714   MORF-related gene X     T154   TGGTGACAGTT   2,00   2,23   Hs.9242   purine-rich element binding protein B     T155   CAGCTCATCTA   2,00   2,23   Hs.285634   Homo sapiens HSPC222 mRNA, complete cds     T156   GGGGTCTGGGC   1,00   1,13   Hs.33026   Human DNA sequence from clone RP3-402G11 on chromoso     T157   ACTGCGAGGAC   1,00   1,13   Hs.110477   dolichyl-phosphate mannosyltransferase polypeptide 3     T158   ACGCAGGCGCC   1,00   1,14   Hs.78103   nucleosome assembly protein 1-like 4     T159   GACTAAGAAAT   1,00   1,14   Hs.66713   hepatitis delta antigen-interacting protein     T160   TGGGCTTGCCT   1,00   1,14   Hs.285163   ESTs     T161   CGCGCCCGGCC   1,00   1,14   Hs.248267   thiosulfate sulfurtransferase (rhodanese)     T162   GTGAAGCTGAT   1,00   1,14   Hs.106671   cleft lip and palate associated transmembrane protei     T163   TGAAACTCATC   2,00   2,27   Hs.85844   neurotrophic tyrosine kinase, receptor,	7148	CTGTGCATTTT	2,00	2,21	Hs.172207	non-POU-domain-containing, octamer-
member 2  7150 GGGTGCTTGGT 1,00 1,12 Hs.6551 ATPase, H+ transporting, lysosomal (vacuolar proton)  7151 AGCTGTGTAAA 1,00 1,12 Hs.279923 putative nucleotide binding protein, estradiol-induc  7152 CAAGATAAATT 1,00 1,12 Hs.177556 melanoma antigen, family D, 1  7153 ATAGAGGCAAT 2,00 2,22 Hs.173714 MORF-related gene X  7154 TGGTGACAGTT 2,00 2,23 Hs.9242 purine-rich element binding protein B  7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds  7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso  7157 ACTGCGAGGAC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4  7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A  7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese)  7162 GTGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,						
Title	7149	AGTTTCCCAAT	3,00	3,27	Hs.75854	
(vacuolar proton 7151 AGCTGTGTAAA 1,00 1,12 Hs.279923 putative nucleotide binding protein, estradiol-induc 7152 CAAGATAAATT 1,00 1,12 Hs.177556 melanoma antigen, family D, 1 7153 ATAGAGGCAAT 2,00 2,22 Hs.173714 MORF-related gene X 7154 TGGTGACAGTT 2,00 2,23 Hs.9242 purine-rich element binding protein B 7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds 7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,						
7151 AGCTGTGTAAA  1,00  1,12 Hs.279923 putative nucleotide binding protein, estradiol-induc  7152 CAAGATAAATT  1,00  1,12 Hs.177556 melanoma antigen, family D, 1  7153 ATAGAGGCAAT  7,00  2,22 Hs.173714 MORF-related gene X  7154 TGGTGACAGTT  2,00  2,23 Hs.9242 purine-rich element binding protein B  7155 CAGCTCATCTA  2,00  2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds  7156 GGGGTCTGGGC  1,00  1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso  7157 ACTGCGAGGAC  1,00  1,14 Hs.78103 nucleosome assembly protein 1-like 4  7159 GACTAAGAAAT  1,00  1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC  1,00  1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC  1,00  1,14 Hs.285163 ESTs  7162 GTGAAGCTGAT  1,00  1,14 Hs.285844 neurotrophic tyrosine kinase, receptor,	7150	GGGTGCTTGGT	1,00	1,12	Hs.6551	
estradiol-induc  7152 CAAGATAAATT 1,00 1,12 Hs.177556 melanoma antigen, family D, 1  7153 ATAGAGGCAAT 2,00 2,22 Hs.173714 MORF-related gene X  7154 TGGTGACAGTT 2,00 2,23 Hs.9242 purine-rich element binding protein B  7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds  7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso  7157 ACTGCGAGGAC 1,00 1,14 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3  7158 ACGCAGGCGCC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4  7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A  7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese)  7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei  7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	- 4 5 1	100000000000000000000000000000000000000				
7152 CAAGATAAATT 1,00 1,12 Hs.177556 melanoma antigen, family D, 1 7153 ATAGAGGCAAT 2,00 2,22 Hs.173714 MORF-related gene X 7154 TGGTGACAGTT 2,00 2,23 Hs.9242 purine-rich element binding protein B 7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds 7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7151	AGCTGTGTAAA	1,00	1,12	Hs.279923	
7153 ATAGAGGCAAT  2,00  2,22 Hs.173714 MORF-related gene X  7154 TGGTGACAGTT  2,00  2,23 Hs.9242 purine-rich element binding protein B  7155 CAGCTCATCTA  2,00  2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds  7156 GGGGTCTGGGC  1,00  1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso  7157 ACTGCGAGGAC  1,00  1,13 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3  7158 ACGCAGGCGCC  1,00  1,14 Hs.78103 nucleosome assembly protein 1-like 4  7159 GACTAAGAAAT  1,00  1,14 Hs.66713 hepatitis delta antigen-interacting protein A  7160 TGGGCTTGCCT  1,00  1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC  1,00  1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese)  7162 GTGAAGCTGAT  1,00  1,14 Hs.106671 cleft lip and palate associated transmembrane protei  7163 TGAAACTCATC  2,00  2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7152	CAAGATAAATT	1,00	1,12	Hs.177556	
7154 TGGTGACAGTT 2,00 2,23 Hs.9242 purine-rich element binding protein B 7155 CAGCTCATCTA 2,00 2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds 7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3- 402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7153	ATAGAGGCAAT				
7155 CAGCTCATCTA  2,00  2,23 Hs.285634 Homo sapiens HSPC222 mRNA, complete cds  7156 GGGGTCTGGGC  1,00  1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso  7157 ACTGCGAGGAC  1,00  1,13 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3  7158 ACGCAGGCGCC  1,00  1,14 Hs.78103 nucleosome assembly protein 1-like 4  7159 GACTAAGAAAT  1,00  1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC  1,00  1,14 Hs.285163 ESTs  7162 GTGAAGCTGAT  1,00  1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese)  7162 GTGAAGCTGAT  1,00  2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7154	TGGTGACAGTT				
7156 GGGGTCTGGGC 1,00 1,13 Hs.33026 Human DNA sequence from clone RP3-402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,13 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3 7158 ACGCAGGCGCC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7155	CAGCTCATCTA	2,00	2,23	Hs.285634	
402G11 on chromoso 7157 ACTGCGAGGAC 1,00 1,13 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3 7158 ACGCAGGCGCC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,						complete cds
7157 ACTGCGAGGAC 1,00 1,13 Hs.110477 dolichyl-phosphate mannosyltransferase polypeptide 3 7158 ACGCAGGCGCC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7156	GGGGTCTGGGC	1,00	1,13	Hs.33026	Human DNA sequence from clone RP3-
7158 ACGCAGGCGCC 1,00 1,14 Hs.78103 nucleosome assembly protein 1-like 4 7159 GACTAAGAAAT 1,00 1,14 Hs.66713 hepatitis delta antigen-interacting protein A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7157	ACTGCGAGGAC	1,00	1,13	Hs.110477	dolichyl-phosphate mannosyltransferase
7159 GACTAAGAAAT  1,00  1,14 Hs.66713 hepatitis delta antigen-interacting protein A  7160 TGGGCTTGCCT  1,00  1,14 Hs.285163 ESTs  7161 CGCGCCCGGCC  1,00  1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese)  7162 GTGAAGCTGAT  1,00  1,14 Hs.106671 cleft lip and palate associated transmembrane protei  7163 TGAAACTCATC  2,00  2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,	7158	ACGCAGGCGCC	1.00	1.14	Hs.78103	
A 7160 TGGGCTTGCCT 1,00 1,14 Hs.285163 ESTs 7161 CGCGCCCGGCC 1,00 1,14 Hs.248267 thiosulfate sulfurtransferase (rhodanese) 7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,						
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7162 GTGAAGCTGAT 1,00 1,14 Hs.106671 cleft lip and palate associated transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,						
transmembrane protei 7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,						
7163 TGAAACTCATC 2,00 2,27 Hs.85844 neurotrophic tyrosine kinase, receptor,			,	,		
	7163	TGAAACTCATC	2,00	2,27	Hs.85844	

	CTGCCAACTTC	2,00			cofilin 1 (non-muscle)
7165	TAGGATGGGGT	1,00	1,15	Hs.76941	ATPase, Na+/K+ transporting, beta 3
					polypeptide
	TACTTGTGTGT	1,00		Hs.6354	stromal cell derived factor receptor 1
7167	TCAGCTTCACC	1,00	1,15	Hs.28707	signal sequence receptor, gamma
					(translocon-associat
7168	CCTCCACCTAG	5,00	5,48	Hs.146354	peroxiredoxin 2
7169	CGGTTACTGTG	3,00	3,39	Hs.49767	NADH dehydrogenase (ubiquinone) Fe-S
					protein 6 (13kD
7170	GAGCCTTGGTG	2,00	2,31	Hs.183994	protein phosphatase 1, catalytic subunit,
					alpha isof
7171	GGAGCTCTGTG	3,00	3,42	Hs.227750	NADH dehydrogenase (ubiquinone) 1
			Ť		beta subcomplex, 4
7172	GGTTTGTGTGT	1,00	1,17	Hs.83954	hypothetical protein
7173	CGATATTCCCC	1,00	1,17	Hs.37616	Human D9 splice variant B mRNA,
		1			complete cds
7174	GGAAGTTCAAA	1,00	1,17	Hs.237536	ESTs, Weakly similar to AF151067_1
		<u> </u>			HSPC233 [H.sapien
7175	AATGGAATGGA	1,00	1,17		ESTs
7176	TCTGCCTGGGG	2,00			MUM2 protein
	AGATCCTACTT	1,00			farnesyl-diphosphate farnesyltransferase
		''-	.,		1
7178	GGAATCCAATC	1,00	1.18	Hs.252587	pituitary tumor-transforming 1
	CTTGCCTGAAG	1,00			bridging integrator 1
	GCCTGGCCATC	2,00			DKFZp564J157 protein
	TCCTGCTGCCG	2,00			KIAA0273 gene product
	CACCTTCCAGC	1,00			melanoma-associated antigen
		.,00		110.27 0000	recognised by cytotoxic
7183	TCCAATACTGC	1,00	1 22	Hs.84153	dynamitin (dynactin complex 50 kD
100	100/11/10/00	1,00	٠,೭೭	113.04100	subunit)
7184	CAGCTGTAGTT	1,00	1 22	Hs.75824	KIAA0174 gene product
	AGAATATCAGT	1,00		Hs.31803	ESTs, Weakly similar to N-WASP
1, 100	NO/MINIONO!	1,00	1,22	113.01000	[H.sapiens]
7186	TCCTAGTAGGA	1,00	1 22	He 252831	reticulon 3
	ACCAAGCTGGG	1,00			tyrosyl-tRNA synthetase
	AAGAATTTGAA	1,00			NADH dehydrogenase (ubiquinone) 1
, 100	700701110701	1,00	1,22	113.100-700	beta subcomplex, 1
7189	GCCCGAGCCC	1,00	1 22	He 178112	DNA segment, single copy probe LNS-
1 103	00000000000	1,00	1,22	113.170112	CAI/LNS-CAII (del
7190	AAGGAAGATCC	1,00	1 22	Hs.11465	glutathione-S-transferase like;
1, 130	/NOO/NO/100	1,00	1,22	113.11400	glutathione transfer
7101	CAGTTTGTACA	1,00	1 22	Hs.1023	pyruvate dehydrogenase (lipoamide)
101	ONOTITION .	1,00	1,22	1 13. 1025	alpha 1
7192	CGTGTTAATGG	3,00	3 60	Hs.2110	zinc finger protein 9 (a cellular retroviral
1 192	OSIGITATIOG	3,00	3,00	1 13.2 ( 10	nucleic
7193	GCGGTTGTGGC	1,00	1 22	Hs.79356	Lysosomal-associated multispanning
1, 193	00001101000	1,00	1,23	1 13.1 3330	membrane protein-
710/	AAAGTTCTCAG	1,00	1 22	He 284242	tetraspan NET-6 protein
	ATACAGAATAA	1,00			delta-like homolog (Drosophila)
1 193	MINUNGANIAN	1,00	1,23	IUS. 109778	Jueita-like nomolog (Drosophila)

7196	ACCTTTACTGT	1,00	1,24	Hs.77356	transferrin receptor (p90, CD71)
	TGGGCCTGGCT	1,00		Hs.6351	cleavage and polyadenylation specific
					factor 4, 30kD
7198	CATAAAGTTTA	1,00	1,24	Hs.3642	DKFZP564B163 protein
7199	TGTCCTGGTTC	2,00	2,47	Hs.179665	cyclin-dependent kinase inhibitor 1A
					(p21, Cip1)
7200	CGGTTTGCATC	1,00	1,25	Hs.31547	NADH dehydrogenase (ubiquinone) 1
					alpha subcomplex,
7201	TTCTGTGTCAC	1,00	1,25	Hs.111024	solute carrier family 25 (mitochondrial carrier; cit
7202	AAGTTCTGCGG	1,00	1.26	Hs.79411	replication protein A2 (32kD)
	TGCTCCTACCC	2,00			Fc fragment of IgG binding protein
$\overline{}$	GTTGCTGCCCT	1,00		Hs.9234	seven transmembrane domain protein
	AGGGTTGGAAG	1,00		Hs.15106	chromosome 14 open reading frame 1
	ATCCGGGGAGC	1,00	<del></del>		putative c-Myc-responsive
	TTTTTGTACAG	2,00		Hs.78040	KDEL (Lys-Asp-Glu-Leu) endoplasmic
					reticulum protein
7208	TTTCAGAGAGA	4,00			signal recognition particle 9kD
	GTTAACGTCCC	5,00		Hs.178391	ribosomal protein L44
	CTGATGGCAGA	1,00		Hs.75874	pregnancy-associated plasma protein A
	CCTTTGGCTAG	1,00			40S ribosomal protein S27 isoform
7212	TAAGTGGAATA	2,00	2,55	Hs.75103	tyrosine 3-monooxygenase/tryptophan 5-
					monooxygenase
7213	GTGCTGGAGAA	3,00	3,79	Hs.53125	small nuclear ribonucleoprotein D2
					polypeptide (16.5
	TAAGGAGCTGA	13,00			ribosomal protein S26
	CACCCCTGATG	7,00			creatine kinase, brain
$\blacksquare$	CCCCTCTGAGT	1,00		Hs.7957	adenosine deaminase, RNA-specific
	ACTACCTTCAC	1,00			px19-like protein
7218	GGGCCCAGGAG	1,00	1,31		ESTs, Weakly similar to diaphanous 1 [H.sapiens]
7219	GGCCCTGAGCG	4,00	5,09		polymerase (RNA) II (DNA directed)
					polypeptide L (7.
	TAACCAATCAG	2,00			RAB5C, member RAS oncogene family
	GCTGCCCTTGA	4,00	5,10	Hs.278242	tubulin, alpha, ubiquitous
7222	TTTGGGGCTGG	2,00	2,62	Hs.7476	ATPase, H+ transporting, lysosomal
					(vacuolar proton
7223	ССССТСССТСС	1,00	1,32	Hs.79410	solute carrier family 4, anion exchanger, member 2 (
7224	AATAAATGGAT	2,00	2.64	Hs.109052	chromosome 14 open reading frame 2
	GGAAAAAAAA	7,00			ATP synthase, H+ transporting,
		,[	., - ,		mitochondrial F1 comp
7226	GGAAAAGTGGT	1,00	1,33	Hs.75621	protease inhibitor 1 (anti-elastase),
					alpha-1-antitr
7227	TGGCTTGCTCA	1,00	1,33	Hs.3688	cisplatin resistance-associated
		·			overexpressed protei
7228	GCCCGTGCCAC	1,00	1,33	Hs.15760	ESTs, Weakly similar to similar to Yeast
					hypothetica
7229	CCACCCCCACC	1,00	1,33	Hs.148101	serum constituent protein

	000000	4 5 5 1		46 ==	1
	GGTGGCTTTGC	1,00			NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 3
7231	AATGCTGGCAA	1,00	1,35	Hs.181195	MRJ gene for a member of the DNAJ protein family
7232	GACCGAGGTGG	1,00	1,35	Hs.129953	Ewing sarcoma breakpoint region 1
7233	CAGAGATGAAT	2,00	2,70	Hs.8997	heat shock 70kD protein 1A
7234	ATCTGAAGCAG	1,00	1,36	Hs.256311	granin-like neuroendocrine peptide precursor
7235	TGCAGCGCCTG	1,00	1,37	Hs.77573	uridine phosphorylase
7236	GCCAGGAAGCC	1,00			hypothetical protein
7237	стттсттств	1,00			polymerase (RNA) II (DNA directed) polypeptide B (14
	GCGCCGCCCCA	1,00	1,37	Hs.108665	zinedin
7239	ATGGGGCAGGG	1,00	1,38	Hs.79226	fasciculation and elongation protein zeta 1 (zygin I
7240	CATTCCTCCTT	1,00	1,38	Hs.2985	emerin (Emery-Dreifuss muscular
					dystrophy)
	ACAACGTCCAG	1,00			p53-responsive gene 2
	CTCTTCGAGAA	3,00			glutathione peroxidase 1
	GGCTCCCACTG	5,00			heat shock 90kD protein 1, beta
7244	GTATCTTCACA	1,00	1,39	Hs.250899	heat shock factor binding protein 1
	CTTAAATCTGG	1,00		Hs.94	heat shock protein, DNAJ-like 2
7246	GATGCGCTTGT	1,00	1,40	Hs.184014	ribosomal protein L31
7247	TAGACTTATTG	1,00	1,40	Hs.170197	glutamic-oxaloacetic transaminase 2, mitochondrial (
7248	ACGATTGATGA	1,00	1,40	Hs.161554	hypothetical protein FLJ20159
7249	GCAACGGGCCC	1,00	1,41	Hs.8679	cytosolic acyl coenzyme A thioester hydrolase
7250	ATAAATTGGGT	1,00	1,41	Hs.81634	ATP synthase, H+ transporting, mitochondrial F0 comp
7251	TTTTAAATTAG	1,00	1,41	Hs.75598	heterogeneous nuclear ribonucleoprotein A2/B1
	CTCGGTGATGT	1,00			Ras homolog enriched in brain 2
	AGATGTGTGGG	1,00			hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl-Coen
	ACGGCTCCGAG	1,00			ESTs
	CCTCTCCTCCC	1,00	1,43	Hs.85004	centromere protein B (80kD)
	TACATTCTGTG	1,00		Hs.86386	myeloid cell leukemia sequence 1 (BCL2-related)
7257	TGCTGACTCCC	1,00	1,45	Hs.29076	ESTs, Highly similar to NEST_HUMAN NESTIN_ [H.sapien
	CCGCTGCTTGT	1,00			heat shock protein, neuronal DNAJ-like 1
	TAACATTAAAG	1,00			X-ray repair complementing defective repair in Chine
	TGGAGCGCTAC	1,00	1,49	Hs.237924	CGI-69 protein
	CCCTCTTTGGA	1,00		Hs.181174	
7262	GTGCTGGACCT	2,00	3,03	Hs.179774	proteasome (prosome, macropain) activator subunit 2
7263	GCCACACCCCA	1,00	1,52	Hs.7718	ESTs, Weakly similar to KIAA1402

					protein [H.sapiens]
7264	TAATTCTTCTC	2,00	3,09	Hs.1708	chaperonin containing TCP1, subunit 3
					(gamma)
7265	GTGTGTGGTGC	1,00	1,56	Hs.151032	Homo sapiens clone 23856 unknown
					mRNA, partial cds
7266	GCAACAGCAAT	3,00	4,60	Hs.9950	Sec61 gamma
7267	GGTGTGGAAGT	1,00	1,57	Hs.9659	hypothetical protein DKFZp434E026
7268	GAGCGGGATGG	2,00		Hs.77060	proteasome (prosome, macropain)
		_,	-,	,	subunit, beta type,
7269	ACACTACGGGT	1,00	1.58	Hs.109494	secreted protein of unknown function
	GCTGGAGCTAG	1,00		Hs.74635	dihydrolipoamide dehydrogenase (E3
		.,	1,00		component of pyru
7271	TAACTTGTGAC	2,00	3 18	Hs 118512	integrin, alpha V (vitronectin receptor,
		2,00	0,10	110.110012	alpha polyp
7272	TAAGATTTCAA	1,00	1.61	Hs.15265	heterogeneous nuclear ribonucleoprotein
1212		1,00	1,01	113.10200	R
7272	ACAAGTACCCA	1,00	1 61	He 142827	P311 protein
	GGGCCCCAAAG	2,00		Hs.256301	
	CATCCTGCTGC			Hs.74619	
1215	CATCCTGCTGC	2,00	3,20	HS.74019	proteasome (prosome, macropain) 26S
7070	00400004044	2.00	2.07	11- 070040	subunit, non-ATP
	CCAGGGGAGAA	2,00			interferon, alpha-inducible protein 27
	TGCTGTGTGCT	1,00		Hs.90606	15 kDa selenoprotein
	TGAAAGTGTGG	1,00		Hs.36927	heat shock 105kD
7279	AAACATTGGGG	1,00	1,66	Hs.8203	endomembrane protein emp70 precursor
					isolog
7280	TGGAACCTTGC	1,00	1,66	Hs.194625	dynein, cytoplasmic, light intermediate
					polypeptide
7281	TGAAGAGAAGA	1,00	1,66	Hs.120	anti-oxidant protein 2 (non-selenium
					glutathione per
7282	CCTGTGACAGC	2,00	-3,32	Hs.120	anti-oxidant protein 2 (non-selenium
					glutathione per
7283	CAACTAATTCA	11,00	17,42	Hs.75106	clusterin (complement lysis inhibitor, SP-
					40,40, sul
7284	AAGGCCTTGTG	1,00	1,67	Hs.74649	cytochrome c oxidase subunit VIc
7285	AGAGCCCTAGG	1,00		Hs.16297	COX17 (yeast) homolog, cytochrome c
		, i			oxidase assembly
7286	TTTTTACTGAT	1,00	1,68	Hs.111577	integral membrane protein 2C
	GCTGGCTGGCT	3,00			chaperonin containing TCP1, subunit 7
		_,	-1- •		(eta)
7288	CGTTCCTGCGG	2,00	3.39	Hs.75424	inhibitor of DNA binding 1, dominant
		_,,55	3,50		negative helix-
7289	CCATTTTTACC	1,00	1 71	Hs.59271	U2(RNU2) small nuclear RNA auxillary
	2	.,55	•,••		factor 1 (non-s
7290	ATGCGGGAGAA	1,00	1 71	Hs 109748	Homo sapiens CAC-1 mRNA, partial cds
	TTTGTGTCACG	1,00		Hs.15093	hypothetical protein
	TCCTCCCTCCC	1,00		Hs.1390	proteasome (prosome, macropain)
1292	10010001000	1,00	1,74	11 15. 1390	1'
7202	GGCACACTAAA	4 00	174	Un 44070	subunit, beta type,
	GGCACAGTAAA	1,00		Hs.11270	ESTs
1294	CTAATAAATGC	1,00	1,/5	Hs.43621	ESTs

7295	TGTCGCTGGGG	2,00	3,49	Hs.227152	mannan-binding lectin serine protease 1 (C4/C2 activ
7296	GGCCATCTCTT	2,00	3,50	Hs.74405	tyrosine 3-monooxygenase/tryptophan 5- monooxygenase
7297	CACTTGAAAAG	1,00	1.76	Hs.7753	calumenin
	GACAGCTGAGC	1,00		Hs.76240	adenylate kinase 1
	GCTGCTCCCTT	1,00		Hs.3804	DKFZP564C1940 protein
	GCGGAGAGAGG	1,00		Hs.286	ribosomal protein L4
	TCAGGCATTTT	2,00		Hs.5566	gap junction protein, beta 2, 26kD (connexin 26)
7302	CAATTAAAAGG	3,00	5.33	Hs 149923	X-box binding protein 1
<del></del>	CAACTTAGTTT	3,00			death-associated protein 6
	GCAGGTGGTTT	1,00			splicing factor (CC1.3)
	GCTGTCATCAG	1,00		Hs.4745	proteasome (prosome, macropain) 26S subunit, ATPase,
7306	GAGCGGGATCA	1,00	1,85	Hs.73737	splicing factor, arginine/serine-rich 1 (splicing fa
7307	ATTAACAAAGC	5,00	9,03	Hs.113368	neuroendocrine secretory protein 55
7308	TGCCGTTTTGA	1,00	1,86	Hs.2006	glutathione S-transferase M3 (brain)
7309	CCGTGCTCATC	2,00	3,72		carbonyl reductase
7310	AAGGCACAGAC	1,00	1,90		CDP-diacylglycerolinositol 3- phosphatidyltransfera
7311	ATTTGTCCCAG	3,00	5,62	Hs.139800	high-mobility group (nonhistone chromosomal) protein
7312	AAGGCCGAGTA	1,00	1,91	Hs.31387	DKFZP564J0123 protein
7313	ATAGTAGCTTC <sub>.</sub>	1,00		Hs.118400	singed (Drosophila)-like (sea urchin fascin homolog
7314	GGACTGGCCCA	1,00	1,93	Hs.263812	nuclear distribution gene C (A.nidulans)
7315	ACTCAGAAGAG	2,00	3,93		NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 2
7316	CTGAGCTGTAC	1,00	1.97	Hs.8737	WD repeat domain 6
	ACTTAAGGAAC	1,00			DKFZP564B167 protein
	GCTATTTGAAA	1,00			seven in absentia (Drosophila) homolog
	TTCTCTCTGTT	1,00	1,99	Hs.77541	ADP-ribosylation factor 5
7320	GGGAGCCCGGG	1,00		Hs.183986	poliovirus receptor-related 2 (herpesvirus entry med
7321	AATATGTGGGC	5,00	9,71		cytochrome c oxidase subunit VIc
7322	GCTCTCTATGC	2,00			signal sequence receptor, delta
					(translocon-associat
7323	TGAGTGGTCAC	1,00	2,01		ESTs, Weakly similar to GEF-2 protein [H.sapiens]
7324	ATTCAGCACCT	1,00	2,04		HSPC033 protein
	CCTGTTCTCCT	1,00			G8 protein
	TGGGAAGTGGG	2,00			hypothetical protein PRO2160
	GATCCCAACTG	5,00			metallothionein 2A
	TTAATAAAAGT	1,00			CGI-113 protein
	CCTCGCTCAGT	1,00			hydroxyacyl-Coenzyme A

7330 ACTCCTGTCCT         1,00         2,11 Hs.7357         DKFZP588N1922 protein           7331 GAGCCGCCTCT         1,00         2,12 Hs.30026         HSPC182 protein           7332 CTTGATTCCCA         1,00         2,13 Hs.77266         quiescin Q6           7333 TGGCAACCTTT         1,00         2,13 Hs.279952         glutathione S-transferase subunit 13 homolog           7334 TTATGGGGAGG         1,00         2,16 Hs.1600         chaperonin containing TCP1, subunit 5 (epsilon)           7335 TCATAGAAACC         1,00         2,17 Hs.82689         tumor rejection antigen (gp96) 1           7337 ACAAACCCCCA         1,00         2,17 Hs.180532 heat shock 90kD protein 1, alpha           7339 ACTGGTACGTG         1,00         2,17 Hs.180532 heat shock 90kD protein 1, alpha           7340 CCAGGAGGAGAT         6,00         12,26 Hs.180414 heat shock 70kd protein 10 (HSC71)           7341 GGGACGAGTGA         3,00         6,45 Hs.337         transmembrane 4 superfamily member 1           7342 GTACCCGGACA         1,00         2,19 Hs.7649         cytochrome c oxidase subunit VIc           7343 GGCTTGGTTTT         1,00         2,22 Hs.172801         Isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,22 Hs.102833 heme oxygenase (decycling) 1           7348 TCCGGGAGGGGT         1,00         2,24 Hs.202833 heme oxygenase		<u> </u>				dahudaganana/2 katanayi Caan		
7331 GAGCGCCTCT         1,00         2,12 Hs.30026         HsPC182 protein           7332 CTTGATTCCCA         1,00         2,13 Hs.77266         quiescin Q6           7333 TGGCAACCTTT         1,00         2,14 Hs.776612         quiescin Q6           7334 TTATGGGGAGG         1,00         2,14 Hs.75612         stress-induced-phosphoprotein 1 (Hsp70/Hsp80-organiz)           7335 TCATAGAAACC         1,00         2,16 Hs.1600         chaperonin containing TCP1, subunit 5 (epsilon)           7338 GAGGAAGAAGA         2,00         4,31 Hs.82689         tumor rejection antigen (gp96) 1           7338 GCTTAGAAGTG         1,00         2,17 Hs.180532         heat shock 90kD protein 1, alpha           7339 ACTGGTACGTG         1,00         2,17 Hs.180532         heat shock 90kD protein 1, alpha           7340 CCAGGAGGAAT         6,00         12,65 Hs.180414         heat shock 70kd protein 10 (HSC71)           7341 GGGACGAGTGA         3,00         6,45 Hs.3337         transmembrane 4 superfamily member 1           7342 GTACCGGACA         1,00         2,19 Hs.74649         eytochrome c oxidase subunit Vol.           7344 TTGGGAGCAGG         1,00         2,29 Hs.7524         coatomer protein complex, subunit beta 2 (beta prime           7345 TCTGCAAAAA         1,00         2,23 Hs.81281         hypothetical protein           7346	7000	AOTOOTOTOOT	4.00	0.44	11. 7057	dehydrogenase/3-ketoacyl-Coen		
7332 CTTGATTCCCA         1,00         2,13 Hs.77266         quiescin Q6           7333 TGGCAACCTTT         1,00         2,13 Hs.279952         glutathione S-transferase subunit 13 homolog           7334 TTATGGGGAGG         1,00         2,14 Hs.75612         stress-induced-phosphoprotein 1 (Hsp70/Hsp80-organiz           7335 TCATAGAAACC         1,00         2,16 Hs.1600         chaperonin containing TCP1, subunit 5 (epsilon)           7336 GAGGAAGAAGA         2,00         4,31 Hs.82689         tumor rejection antigen (gp96) 1           7337 ACAAACCCCCA         1,00         2,17 Hs.186239         hat Fase, Na+/K+ transporting, beta 1 polypeptide           7338 GCTTAGAAGTG         1,00         2,17 Hs.186529         hat shock 90kD protein 1, alpha 1 polypeptide           7339 ACTGGTACGTG         1,00         2,17 Hs.1865751         ATP say, and the transporting, mitochondrial F0 comp mitoc								
7333         TGGCAACCTTT         1,00         2,13         Hs.279952 glutathione S-transferase subunit 13 homolog           7334         TTATGGGGAGG         1,00         2,14         Hs.75612 stress-induced-phosphoprotein 1 (Hsp70/Hsp90-organiz)           7335         TCATAGAAACC         1,00         2,16         Hs.1600 chaperonin containing TCP1, subunit 5 (epsilon)           7336         GAGGAAGAGA         2,00         4,31         Hs.82689 lumor rejection antigen (gp96) 1           7337         ACAAACCCCCA         1,00         2,17         Hs.180532 heat shock 90kD protein 1, alpha           7338         GCTTAGAAGTG         1,00         2,17         Hs.180532 heat shock 90kD protein 1, alpha           7339         ACTGGTACGTG         1,00         2,17         Hs.180532 heat shock 90kD protein 1, alpha           7340         CCAGGAGGAAT         6,00         12,65         Hs.180414 heat shock 70kd protein 10 (HSC71)           7341         GGGACGAGTGA         1,00         2,19         Hs.75724 coatomer protein complex, subunit beta           7342         GTCCGGACA         1,00         2,22         Hs.172801 isoleucine-tRNA synthetase           7345         TCTGCAAAAAA         1,00         2,23         Hs.19303 heme oxygenase (decycling) 1           7346         GGGGGTGGGG         1,00 <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td></td<>								
Name								
(Hsp70/Hsp90-organiz chaperonin containing TCP1, subunit 5 (epsilon)	7333	TGGCAACCTTT	1,00		·	homolog		
TCATAGAAACC	7334	TTATGGGGAGG	1,00	2,14	Hs.75612			
7336 GAGGAAGAAGA         2,00         4,31 Hs.82689         tumor rejection antigen (gp96) 1           7337 ACAAACCCCCA         1,00         2,17 Hs.78629         ATPase, Na+/K+ transporting, beta 1 polypeptide           7338 GCTTAGAAGTG         1,00         2,17 Hs.180532 heat shock 90kD protein 1, alpha           7339 ACTGGTACGTG         1,00         2,17 Hs.155751 ATP synthase, H+ transporting, mitochondrial F0 comp           7340 CCAGGAGGAAT         6,00         12,65 Hs.180414 heat shock 70kd protein 10 (HSC71)           7341 GGGACGAGTGA         3,00         6,45 Hs.3337         transmembrane 4 superfamily member 1           7342 GTACCCGGACA         1,00         2,19 Hs.74649         cytochrome c oxidase subunit VIc           7343 GGCTTTGATTT         1,00         2,20 Hs.75724         coatomer protein complex, subunit beta 2 (beta prime           7344 TTGGGAGCAGG         1,00         2,22 Hs.172801         isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,23 Hs.91281         hypothetical protein           7346 GGGGTGGGG         1,00         2,27 Hs.202833         heme oxygenase (decycling) 1           7347 CGTGGGTGGCC         1,00         2,27 Hs.202833         heme oxygenase (decycling) 1           7349 GAGAACCGTAG         2,00         4,57 Hs.105547 neural proliferation, differentiation and control, 1         neural proliferation,	7335	TCATAGAAACC	1,00	2,16	Hs.1600	chaperonin containing TCP1, subunit 5		
7337   ACAAACCCCCA   1,00   2,17   Hs.78629   ATPase, Na+/K+ transporting, beta 1 polypeptide   1,00   2,17   Hs.180532   heat shock 90kD protein 1, alpha   1,00   2,17   Hs.180532   heat shock 90kD protein 1, alpha   1,00   2,17   Hs.185751   ATP synthase, H+ transporting, mitochondrial F0 comp   1,00   2,17   Hs.180414   heat shock 70kd protein 10 (HSC71)   1,00   2,19   Hs.74649   cytochrome c oxidase subunit VIc   2,30   Hs.75724   Coatomer protein complex, subunit beta   2 (beta prime   2 (beta pri	7336	GAGGAAGAAGA	2,00	4,31	Hs.82689			
7338 GCTTAGAAGTG         1,00         2,17 Hs.180532 heat shock 90kD protein 1, alpha           7339 ACTGGTACGTG         1,00         2,17 Hs.155751 ATP synthase, H+ transporting, mitochondrial F0 comp           7340 CCAGGAGGAAT         6,00         12,65 Hs.180414 heat shock 70kd protein 10 (HSC71)           7341 GGGACGAGTGA         3,00         6,45 Hs.3337 transmembrane 4 superfamily member 1           7342 GTACCCGGACA         1,00         2,19 Hs.74649 cytochrome c oxidase subunit VIc coatomer protein complex, subunit beta 2 (beta prime           7343 GGCTTTGATTT         1,00         2,22 Hs.172801 isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,22 Hs.172801 isoleucine-tRNA synthetase           7346 GGGGATGGGGT         1,00         2,25 Hs.99093 Homo sapiens chromosome 19, cosmid R28379           7347 CGTGGGTGGGG         1,00         2,27 Hs.202833 heme oxygenase (decycling) 1           7348 TTCCGCGTGCC         1,00         2,29 Hs.153357 procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349 GAGAACCGTAG         2,00         4,57 Hs.105547           7350 TTACGAGGAAG         1,00         2,32 Hs.227949 SEC13 (S. cerevisiae)-like 1           7351 TTGCTTTTGTT         1,00         2,33 Hs.75290           7352 GAAGATGTGTG         3,00         7,05 Hs.112318 6.2 kd protein           7353 GATTTTGTAGC         1,00         2,39 Hs.						ATPase, Na+/K+ transporting, beta 1		
7339 ACTGGTACGTG         1,00         2,17 Hs.155751 ATP synthase, H+ transporting, mitochondrial F0 comp           7340 CCAGGAGGAAT         6,00         12,65 Hs.180414 heat shock 70kd protein 10 (HSC71)           7341 GGGACGAGTGA         3,00         6,45 Hs.3337 transmembrane 4 superfamily member 1           7342 GTACCCGGACA         1,00         2,19 Hs.74649 cytochrome c oxidase subunit VIc           7343 GGCTTTGATTT         1,00         2,20 Hs.75724 coatomer protein complex, subunit beta 2 (beta prime           7344 TTGGGAGCAGG         1,00         2,22 Hs.172801 isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,23 Hs.81281 hypothetical protein           7346 GGGGTGGGGG         1,00         2,27 Hs.202833 heme oxygenase (decycling) 1           7347 CGTGGGTGGGG         1,00         2,27 Hs.202833 heme oxygenase (decycling) 1           7348 TTCCGCGTGCC         1,00         2,29 Hs.153357 procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349 GAGAACCGTAG         2,00         4,57 Hs.105547 neural proliferation, differentiation and control, 1           7350 TTACGAGGAAG         1,00         2,32 Hs.227949 SEC13 (S. cerevisiae)-like 1           7351 TTGCTTTTGTT         1,00         2,33 Hs.75290 ADP-ribosylation factor 4           7352 GAGGTTAGC         1,00         2,39 Hs.84264 acidic protein rich in leucines           7354 TCGTCGCAGAA	7220	CCTTACAACTC	1.00	2 47	Un 100522			
mitochondrial F0 comp   mitochondrial F0 condition   mitochondrial F0 comp   mitochondrial F0 condition								
7341 GGGACGAGTGA         3,00         6,45 Hs.3337         transmembrane 4 superfamily member 1           7342 GTACCCGGACA         1,00         2,19 Hs.74649         cytochrome c oxidase subunit VIc           7343 GCTTTGATTT         1,00         2,20 Hs.75724         coatomer protein complex, subunit beta 2 (beta prime           7344 TTGGGAGCAGG         1,00         2,22 Hs.172801         isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,23 Hs.81281         hypothetical protein           7346 GGGGATGGGGT         1,00         2,25 Hs.99093         Homo sapiens chromosome 19, cosmid R28379           7347 CGTGGGTGGGG         1,00         2,27 Hs.202833 heme oxygenase (decycling) 1           7348 TTCCGCGTGCC         1,00         2,29 Hs.153357           7349 GAGAACCGTAG         2,00         4,57 Hs.105547           7350 TTACGAGGAAG         1,00         2,32 Hs.227949         SEC13 (S. cerevisiae)-like 1           7351 TTGCTTTTGTT         1,00         2,33 Hs.75290         ADP-ribosylation factor 4           7352 GAAGATGTGTG         3,00         7,05 Hs.112318 6.2 kd protein           7353 GATTTTGTAGC         1,00         2,39 Hs.84264         acidic protein rich in leucines           7355 GAGGCTCAATC         1,00         2,40 Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)			1,00			mitochondrial F0 comp		
7342 GTACCCGGACA         1,00         2,19 Hs.74649         cytochrome c oxidase subunit VIc           7343 GGCTTTGATTT         1,00         2,20 Hs.75724         coatomer protein complex, subunit beta           7344 TTGGGAGCAGG         1,00         2,22 Hs.172801         isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,23 Hs.81281         hypothetical protein           7346 GGGATGGGGT         1,00         2,25 Hs.99093         Homo sapiens chromosome 19, cosmid R28379           7347 CGTGGGTGGGG         1,00         2,27 Hs.202833         heme oxygenase (decycling) 1           7348 TTCCGCGTGCC         1,00         2,29 Hs.153357         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349 GAGAACCGTAG         2,00         4,57 Hs.105547         neural proliferation, differentiation and control, 1           7350 TTACGAGGAAG         1,00         2,32 Hs.227949         SEC13 (S. cerevisiae)-like 1           7351 TTGCTTTGTT         1,00         2,33 Hs.75290         ADP-ribosylation factor 4           7352 GAGATGTGTG         3,00         7,05 Hs.112318 6.2 kd protein           7353 GATTTTGTAGC         1,00         2,39 Hs.84264         acidic protein rich in leucines           7354 TCGTCGCAGAA         1,00         2,40 Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma) <t< td=""><td>7340</td><td>CCAGGAGGAAT</td><td></td><td></td><td></td><td></td></t<>	7340	CCAGGAGGAAT						
1,00	7341	GGGACGAGTGA	3,00			transmembrane 4 superfamily member 1		
2 (beta prime     2 (beta prime     7344 TTGGGAGCAGG   1,00   2,22 Hs.172801   isoleucine-tRNA synthetase   7345 TCTGCAAAAAA   1,00   2,23 Hs.81281   hypothetical protein   7346 GGGGATGGGGT   1,00   2,25 Hs.99093   Homo sapiens chromosome 19, cosmid R28379   7347 CGTGGGTGGGG   1,00   2,27 Hs.202833   heme oxygenase (decycling) 1   7348 TTCCGCGTGCC   1,00   2,29 Hs.153357   procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v   7349 GAGAACCGTAG   2,00   4,57 Hs.105547   neural proliferation, differentiation and control, 1   7350 TTACGAGGAAG   1,00   2,32 Hs.227949   SEC13 (S. cerevisiae)-like 1   7351 TTGCTTTTGTT   1,00   2,33 Hs.75290   ADP-ribosylation factor 4   7352 GAAGATGTGTG   3,00   7,05 Hs.112318   6.2 kd protein   7353 GATTTTGTAGC   1,00   2,39 Hs.84264   acidic protein rich in leucines   7354 TCGTCGCAGAA   1,00   2,39 Hs.19561   NADH dehydrogenase (ubiquinone) 1   alpha subcomplex,   chromobox homolog 3 (Drosophila HP1 gamma)   7356 TAGAAAAATAA   1,00   2,42 Hs.944   glucose phosphate isomerase   7357 CCGGGTGATGG   2,00   4,84 Hs.279910   ATX1 (antioxidant protein 1, yeast)   homolog 1   7358 GAAGCTTTGCA   4,00   9,54 Hs.180532   heat shock 90kD protein 1, alpha   7359 GGTTATTTTGG   1,00   2,47 Hs.82085   plasminogen activator inhibitor, type   7360 TGTCGCGGG   2,00   4,97 Hs.76847   KIAA0088 protein   7361 ATGGCGATCTA   1,00   2,56 Hs.73722   APEX nuclease (multifunctional DNA repair enzyme)   7363 GGTGCAGAGCC   1,00   2,56 Hs.539   ribosomal protein S29   10   10   10   10   10   10   10   1	7342	GTACCCGGACA	1,00	2,19	Hs.74649	cytochrome c oxidase subunit VIc		
7344 TTGGGAGCAGG         1,00         2,22 Hs.172801 isoleucine-tRNA synthetase           7345 TCTGCAAAAAA         1,00         2,23 Hs.81281 hypothetical protein           7346 GGGGATGGGGT         1,00         2,25 Hs.99093 Homo sapiens chromosome 19, cosmid R28379           7347 CGTGGGTGGG         1,00         2,27 Hs.202833 heme oxygenase (decycling) 1           7348 TCCGCGTGCC         1,00         2,29 Hs.153357 procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349 GAGAACCGTAG         2,00         4,57 Hs.105547 hs.105547 neural proliferation, differentiation and control, 1           7350 TTACGAGGAAG         1,00         2,32 Hs.227949 SEC13 (S. cerevisiae)-like 1           7351 TTGCTTTTGTT         1,00         2,33 Hs.75290 ADP-ribosylation factor 4           7352 GAGATGTGTG         3,00         7,05 Hs.112318 6.2 kd protein           7353 GATTTGTAGC         1,00         2,39 Hs.84264 acidic protein rich in leucines           7354 TCGTCGCAGAA         1,00         2,39 Hs.19561 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, chromobox homolog 3 (Drosophila HP1 gamma)           7355 GAGGCTCAATC         1,00         2,40 Hs.8123 chromobox homolog 3 (Drosophila HP1 gamma)           7356 TAGAAAAATAA         1,00         2,42 Hs.944 glucose phosphate isomerase           7357 CCGGGTGATGG         2,00 4,84 Hs.279910 ATX1 (antioxidant protein 1, yeast) homolog 1           7358 GAA	7343	GGCTTTGATTT	1,00	2,20	Hs.75724	coatomer protein complex, subunit beta		
7345 TCTGCAAAAAA         1,00         2,23 Hs.81281         hypothetical protein           7346 GGGGATGGGGT         1,00         2,25 Hs.99093         Homo sapiens chromosome 19, cosmid R28379           7347 CGTGGGTGGG         1,00         2,27 Hs.202833         heme oxygenase (decycling) 1           7348 TCCGCGTGCC         1,00         2,29 Hs.153357         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349 GAGAACCGTAG         2,00         4,57 Hs.105547         neural proliferation, differentiation and control, 1           7350 TTACGAGGAAG         1,00         2,32 Hs.227949         SEC13 (S. cerevisiae)-like 1           7351 TTGCTTTTGTT         1,00         2,33 Hs.75290         ADP-ribosylation factor 4           7352 GAAGATGTGTG         3,00         7,05 Hs.112318 6.2 kd protein           7354 TCGTCGCAGAA         1,00         2,39 Hs.9561         NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, chromobox homolog 3 (Drosophila HP1 gamma)           7355 GAGGCTCAATC         1,00         2,40 Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)           7356 TAGAAAAATAA         1,00         2,42 Hs.944         glucose phosphate isomerase           7357 CCGGGTGATGG         2,00         4,84 Hs.279910         ATX1 (antioxidant protein 1, alpha           7359 GAGAGCTTTGCA         4,00         9,54 Hs.180532         heat sh								
7346         GGGGATGGGGT         1,00         2,25         Hs. 99093         Homo sapiens chromosome 19, cosmid R28379           7347         CGTGGGTGGG         1,00         2,27         Hs. 202833         heme oxygenase (decycling) 1           7348         TTCCGCGTGCC         1,00         2,29         Hs. 153357         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349         GAGAACCGTAG         2,00         4,57         Hs. 105547         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7350         TTACGAGGAAG         1,00         2,32         Hs. 227949         SEC13 (S. cerevisiae)-like 1           7351         TTGCTTTTGTT         1,00         2,33         Hs. 75290         ADP-ribosylation factor 4           7352         GAGATGTGTG         3,00         7,05         Hs. 112318 6.2 kd protein           7353         GATTTGTAGC         1,00         2,39         Hs. 84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs. 8123         chromobox homolog 3 (Drosophila HP1 gamma)           7355         GAGGCTCAATC         1,00         2,40         Hs. 8123         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs. 279910	7344	TTGGGAGCAGG	1,00	2,22	Hs.172801	isoleucine-tRNA synthetase		
7346         GGGGATGGGGT         1,00         2,25         Hs. 99093         Homo sapiens chromosome 19, cosmid R28379           7347         CGTGGGTGGG         1,00         2,27         Hs. 202833         heme oxygenase (decycling) 1           7348         TTCCGCGTGCC         1,00         2,29         Hs. 153357         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349         GAGAACCGTAG         2,00         4,57         Hs. 105547         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7350         TTACGAGGAAG         1,00         2,32         Hs. 227949         SEC13 (S. cerevisiae)-like 1           7351         TTGCTTTTGTT         1,00         2,33         Hs. 75290         ADP-ribosylation factor 4           7352         GAGATGTGTG         3,00         7,05         Hs. 112318 6.2 kd protein           7353         GATTTGTAGC         1,00         2,39         Hs. 84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs. 8123         chromobox homolog 3 (Drosophila HP1 gamma)           7355         GAGGCTCAATC         1,00         2,40         Hs. 8123         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs. 279910	7345	TCTGCAAAAAA	1,00	2,23	Hs.81281	hypothetical protein		
7347         CGTGGGTGGGG         1,00         2,27         Hs.202833         heme oxygenase (decycling) 1           7348         TTCCGCGTGCC         1,00         2,29         Hs.153357         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349         GAGAACCGTAG         2,00         4,57         Hs.105547         neural proliferation, differentiation and control, 1           7350         TTACGAGGAAG         1,00         2,32         Hs.227949         SEC13 (S. cerevisiae)-like 1           7351         TTGCTTTTGTT         1,00         2,33         Hs.75290         ADP-ribosylation factor 4           7352         GAAGATGTGG         3,00         7,05         Hs.112318 6.2 kd protein           7353         GATTTTGTAGC         1,00         2,39         Hs.84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs.19561         NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,           7355         GAGGCTCAATC         1,00         2,40         Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)           7356         TAGAAAAAAAAAAAA         1,00         2,42         Hs.944         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs.180532 </td <td>7346</td> <td>GGGGATGGGGT</td> <td></td> <td>2,25</td> <td>Hs.99093</td> <td></td>	7346	GGGGATGGGGT		2,25	Hs.99093			
7348         TTCCGCGTGCC         1,00         2,29         Hs.153357         procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3v           7349         GAGAACCGTAG         2,00         4,57         Hs.105547         neural proliferation, differentiation and control, 1           7350         TTACGAGGAAG         1,00         2,32         Hs.227949         SEC13 (S. cerevisiae)-like 1           7351         TTGCTTTTGTT         1,00         2,33         Hs.75290         ADP-ribosylation factor 4           7352         GAAGATGTGTG         3,00         7,05         Hs.112318         6.2 kd protein           7353         GATTTTGTAGC         1,00         2,39         Hs.84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs.19561         NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,           7355         GAGGCTCAATC         1,00         2,40         Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)           7356         TAGAAAAATAA         1,00         2,42         Hs.944         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs.279910         ATX1 (antioxidant protein 1, yeast) homolog 1           7358         GAAGCTTTGCA         4,00         9,	7347	CGTGGGTGGGG	1,00	2,27	Hs.202833	heme oxygenase (decycling) 1		
7349         GAGAACCGTAG         2,00         4,57         Hs.105547         neural proliferation, differentiation and control, 1           7350         TTACGAGGAAG         1,00         2,32         Hs.227949         SEC13 (S. cerevisiae)-like 1           7351         TTGCTTTTGTT         1,00         2,33         Hs.75290         ADP-ribosylation factor 4           7352         GAAGATGTGTG         3,00         7,05         Hs.112318         6.2 kd protein           7353         GATTTTGTAGC         1,00         2,39         Hs.84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs.19561         NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,           7355         GAGGCTCAATC         1,00         2,40         Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)           7356         TAGAAAAAATAA         1,00         2,42         Hs.944         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs.279910         ATX1 (antioxidant protein 1, yeast) homolog 1           7358         GAAGCTTTGCA         4,00         9,54         Hs.180532         heat shock 90kD protein 1, alpha           7359         GGTTATTTTGG         1,00         2,47 <t< td=""><td>7348</td><td>TTCCGCGTGCC</td><td></td><td>2,29</td><td>Hs.153357</td><td>procollagen-lysine, 2-oxoglutarate 5-</td></t<>	7348	TTCCGCGTGCC		2,29	Hs.153357	procollagen-lysine, 2-oxoglutarate 5-		
7350         TTACGAGGAAG         1,00         2,32         Hs.227949         SEC13 (S. cerevisiae)-like 1           7351         TTGCTTTTGTT         1,00         2,33         Hs.75290         ADP-ribosylation factor 4           7352         GAAGATGTGTG         3,00         7,05         Hs.112318         6.2 kd protein           7353         GATTTTGTAGC         1,00         2,39         Hs.84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs.19561         NADH dehydrogenase (ubiquinone) 1           7355         GAGGCTCAATC         1,00         2,40         Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)           7356         TAGAAAAATAA         1,00         2,42         Hs.944         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs.279910         ATX1 (antioxidant protein 1, yeast) homolog 1           7358         GAAGCTTTGCA         4,00         9,54         Hs.180532         heat shock 90kD protein 1, alpha           7359         GGTTATTTTGG         1,00         2,47         Hs.76847         KIAA0088 protein           7361         ATGGCGATCTA         1,00         2,53         Hs.73722         APEX nuclease (multifunction	7349	GAGAACCGTAG	2,00	4,57	Hs.105547	neural proliferation, differentiation and		
7351         TTGCTTTTGTT         1,00         2,33         Hs.75290         ADP-ribosylation factor 4           7352         GAAGATGTGTG         3,00         7,05         Hs.112318         6.2 kd protein           7353         GATTTTGTAGC         1,00         2,39         Hs.84264         acidic protein rich in leucines           7354         TCGTCGCAGAA         1,00         2,39         Hs.19561         NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,           7355         GAGGCTCAATC         1,00         2,40         Hs.8123         chromobox homolog 3 (Drosophila HP1 gamma)           7356         TAGAAAAATAA         1,00         2,42         Hs.944         glucose phosphate isomerase           7357         CCGGGTGATGG         2,00         4,84         Hs.279910         ATX1 (antioxidant protein 1, yeast) homolog 1           7358         GAAGCTTTGCA         4,00         9,54         Hs.180532 heat shock 90kD protein 1, alpha           7359         GGTTATTTTGG         1,00         2,47         Hs.82085 plasminogen activator inhibitor, type I           7360         TGTGCTCGGGG         2,00         4,97         Hs.76847 KIAA0088 protein           7361         ATGGCGATCTA         1,00         2,56         Hs.73722 APEX nuclease (multifunctional DNA repair enzyme)	7350	TTACGAGGAAG	1,00	2,32	Hs.227949			
7352 GAAGATGTGTG         3,00         7,05 Hs.112318 6.2 kd protein           7353 GATTTTGTAGC         1,00         2,39 Hs.84264 acidic protein rich in leucines           7354 TCGTCGCAGAA         1,00         2,39 Hs.19561 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,           7355 GAGGCTCAATC         1,00         2,40 Hs.8123 chromobox homolog 3 (Drosophila HP1 gamma)           7356 TAGAAAAATAA         1,00         2,42 Hs.944 glucose phosphate isomerase           7357 CCGGGTGATGG         2,00 4,84 Hs.279910 ATX1 (antioxidant protein 1, yeast) homolog 1           7358 GAAGCTTTGCA         4,00 9,54 Hs.180532 heat shock 90kD protein 1, alpha           7359 GGTTATTTTGG         1,00 2,47 Hs.82085 plasminogen activator inhibitor, type I           7360 TGTGCTCGGGG         2,00 4,97 Hs.76847 KIAA0088 protein           7361 ATGGCGATCTA         1,00 2,53 Hs.180450 ribosomal protein S24           7362 AAAATAAAGAG         1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme)           7363 GGTGCAGAGCC         1,00 2,56 Hs.539 ribosomal protein S29								
7353 GATTTTGTAGC 7354 TCGTCGCAGAA 1,00 2,39 Hs.19561 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 7355 GAGGCTCAATC 1,00 2,40 Hs.8123 chromobox homolog 3 (Drosophila HP1 gamma) 7356 TAGAAAAATAA 1,00 2,42 Hs.944 glucose phosphate isomerase 7357 CCGGGTGATGG 2,00 4,84 Hs.279910 ATX1 (antioxidant protein 1, yeast) homolog 1 7358 GAAGCTTTGCA 4,00 9,54 Hs.180532 heat shock 90kD protein 1, alpha 7359 GGTTATTTTGG 1,00 2,47 Hs.82085 plasminogen activator inhibitor, type I 7360 TGTGCTCGGGG 2,00 4,97 Hs.76847 KIAA0088 protein 7361 ATGGCGATCTA 1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme) 7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29								
TCGTCGCAGAA 1,00 2,39 Hs.19561 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex,  7355 GAGGCTCAATC 1,00 2,40 Hs.8123 chromobox homolog 3 (Drosophila HP1 gamma)  7356 TAGAAAAATAA 1,00 2,42 Hs.944 glucose phosphate isomerase  7357 CCGGGTGATGG 2,00 4,84 Hs.279910 ATX1 (antioxidant protein 1, yeast) homolog 1  7358 GAAGCTTTGCA 4,00 9,54 Hs.180532 heat shock 90kD protein 1, alpha  7359 GGTTATTTTGG 1,00 2,47 Hs.82085 plasminogen activator inhibitor, type I  7360 TGTGCTCGGGG 2,00 4,97 Hs.76847 KIAA0088 protein  7361 ATGGCGATCTA 1,00 2,53 Hs.180450 ribosomal protein S24  7362 AAAATAAAGAG 1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme)  7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29								
7355 GAGGCTCAATC 1,00 2,40 Hs.8123 chromobox homolog 3 (Drosophila HP1 gamma)  7356 TAGAAAAATAA 1,00 2,42 Hs.944 glucose phosphate isomerase  7357 CCGGGTGATGG 2,00 4,84 Hs.279910 ATX1 (antioxidant protein 1, yeast) homolog 1  7358 GAAGCTTTGCA 4,00 9,54 Hs.180532 heat shock 90kD protein 1, alpha  7359 GGTTATTTTGG 1,00 2,47 Hs.82085 plasminogen activator inhibitor, type I  7360 TGTGCTCGGGG 2,00 4,97 Hs.76847 KIAA0088 protein  7361 ATGGCGATCTA 1,00 2,53 Hs.180450 ribosomal protein S24  7362 AAAATAAAGAG 1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme)  7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29						NADH dehydrogenase (ubiquinone) 1		
7356 TAGAAAAATAA1,002,42 Hs.944glucose phosphate isomerase7357 CCGGGTGATGG2,004,84 Hs.279910ATX1 (antioxidant protein 1, yeast) homolog 17358 GAAGCTTTGCA4,009,54 Hs.180532 heat shock 90kD protein 1, alpha7359 GGTTATTTTGG1,002,47 Hs.82085 plasminogen activator inhibitor, type I7360 TGTGCTCGGGG2,004,97 Hs.76847 KIAA0088 protein7361 ATGGCGATCTA1,002,53 Hs.180450 ribosomal protein S247362 AAAATAAAGAG1,002,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme)7363 GGTGCAGAGCC1,002,56 Hs.539 ribosomal protein S29	7355	GAGGCTCAATC	1,00	2,40	Hs.8123	chromobox homolog 3 (Drosophila HP1		
7357 CCGGGTGATGG 2,00 4,84 Hs.279910 ATX1 (antioxidant protein 1, yeast) homolog 1 7358 GAAGCTTTGCA 4,00 9,54 Hs.180532 heat shock 90kD protein 1, alpha 7359 GGTTATTTTGG 1,00 2,47 Hs.82085 plasminogen activator inhibitor, type I 7360 TGTGCTCGGGG 2,00 4,97 Hs.76847 KIAA0088 protein 7361 ATGGCGATCTA 1,00 2,53 Hs.180450 ribosomal protein S24 7362 AAAATAAAGAG 1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme) 7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29	7356	TAGAAAAATAA	1.00	2.42	Hs.944			
homolog 1   7358 GAAGCTTTGCA								
7359 GGTTATTTTGG1,002,47 Hs.82085plasminogen activator inhibitor, type I7360 TGTGCTCGGGG2,004,97 Hs.76847KIAA0088 protein7361 ATGGCGATCTA1,002,53 Hs.180450 ribosomal protein S247362 AAAATAAAGAG1,002,56 Hs.73722APEX nuclease (multifunctional DNA repair enzyme)7363 GGTGCAGAGCC1,002,56 Hs.539ribosomal protein S29	· ·					homolog 1		
7360 TGTGCTCGGGG         2,00         4,97 Hs.76847         KIAA0088 protein           7361 ATGGCGATCTA         1,00         2,53 Hs.180450 ribosomal protein S24           7362 AAAATAAAGAG         1,00         2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme)           7363 GGTGCAGAGCC         1,00         2,56 Hs.539 ribosomal protein S29								
7361 ATGGCGATCTA 1,00 2,53 Hs.180450 ribosomal protein S24 7362 AAAATAAAGAG 1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme) 7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29				<u>-</u>				
7362 AAAATAAAGAG 1,00 2,56 Hs.73722 APEX nuclease (multifunctional DNA repair enzyme) 7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29								
repair enzyme) 7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29			1,00					
7363 GGTGCAGAGCC 1,00 2,56 Hs.539 ribosomal protein S29	7362	AAAATAAAGAG	1,00	2,56	Hs.73722			
	7363	GGTGCAGAGCC	1.00	2.56	Hs.539			
	<del></del>		1,00			myeloid leukemia factor 2		

7365	TGACTGAAGCC	1,00	2,62	Hs.3343	phosphoglycerate dehydrogenase		
	AACGCGGCCAA	7,00		Hs.73798	macrophage migration inhibitory factor		
L					(glycosylatio		
7367	GCCGTGTAGAC	1,00	2,69	Hs.83384	S100 calcium-binding protein, beta		
					(neural)		
7368	AGGGTGAAACT	1,00		Hs.77608	splicing factor, arginine/serine-rich 9		
7369	AAGGTAATGCT	1,00		Hs.23990	hypothetical protein FLJ20479		
7370	ACAGTGGGGAT	2,00	5,54	Hs.75839	zinc finger protein 6 (CMPX1)		
7371	GAACCCTGGGA	1,00	2,78	Hs.155637	protein kinase, DNA-activated, catalytic polypeptide		
7372	TAAATAATTTC	1,00	2,79	Hs.1197	heat shock 10kD protein 1 (chaperonin 10)		
7373	GAAATAAAAGC	1,00	2,81	Hs.57783	eukaryotic translation initiation factor 3, subunit		
7374	GAATTAACATT	2,00	5,70	Hs.79474	tyrosine 3-monooxygenase/tryptophan 5-monooxygenase		
7375	GGTAGCAGGGA	1,00	2,88	Hs.179817	CGI-82 protein		
7376	GTTTAAATCGA	1,00			proteasome (prosome, macropain) subunit, alpha type,		
7377	GACGTCTTAAT	1,00	3,01	Hs.251531	1 proteasome (prosome, macropain) subunit, alpha type,		
7378	TACTAGTCCTC	2,00	6,04	Hs.180532	heat shock 90kD protein 1, alpha		
7379	CGATTCTGGAG	1,00			hypothetical protein		
7380	ACTGGTAAAAA	1,00			ATP synthase, H+ transporting, mitochondrial F0 comp		
7381	AAGGAGTTTGA	1,00	3,05	Hs.661	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 7		
7382	GTGACAACACT	1,00	3.06	Hs.149155	voltage-dependent anion channel 1		
	GGAGTCATTGT	1,00		Hs.82793	proteasome (prosome, macropain) subunit, beta type,		
7384	TGTATAAAAAT	1,00	3.12	Hs.82689	tumor rejection antigen (gp96) 1		
$\overline{}$	GCAGACATTGA	1,00			nucleosome assembly protein 1-like 1		
	GAGGATGGTGT	1,00		Hs.19555	prostate tumor over expressed gene 1		
	TCAAATGCATC	1,00		Hs.182447			
7388	TGCTTGTCCCT.	1,00	3,35	Hs.74571	ADP-ribosylation factor 1		
	AATGTGAGTCA	1,00		Hs.25001	tyrosine 3-monooxygenase/tryptophan 5- monooxygenase		
7390	CGGCCCAACGC	1,00	3,41	Hs.20521	HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 2		
7391	GTTTCATCTCC	1,00	3.42	Hs.13451	dystrobrevin, beta		
	AAGGAATCGGG	1,00		Hs.89545	proteasome (prosome, macropain)		
		,	•		subunit, beta type,		
7393	AAAGTGAAGAT	1,00	3,44	Hs.284158	Homo sapiens mRNA; cDNA DKFZp762B195 (from clone DKF		
7394	TCAGAAGGTGC	1,00	3.49	Hs.74649	cytochrome c oxidase subunit VIc		
	GGGCCTGTGCC	1,00		Hs.85838	solute carrier family 16 (monocarboxylic acid transp		
7396	AGAAATACCAA	1,00	3 56	Hs.74649	cytochrome c oxidase subunit VIc		
1,000	, NOCKY I ACCES	1,00	2,30	1 13.1 4043	Cytochionie c Oxidase Subutilit VIC		

7307	GGAAGTTTCGA	1,00	3 56	Hs.55847	hypothetical protein
	GGGGGTAACTA	1,00			fusion, derived from t(12;16) malignant
1, 290	GGGGGTAACIA	1,00	3,00	10.00000	liposarcoma
7399	CATTTCATAAC	1,00	3.68	Hs.73851	ATP synthase, H+ transporting,
		',55	5,00		mitochondrial F0 comp
7400	GGCTTTGGAGT	1,00	3,73	Hs.90918	chromosome 11 open reading frame 10
7401	GGCGTCCTGGC	1,00		Hs.44017	SIR2 (silent mating type information
			·		regulation 2, S
	TAGACTAGCAA	1,00			tetraspan 3
7403	CCGATCACCGG	1,00	4,18	Hs.12163	eukaryotic translation initiation factor 2,
7404	CAATOCAACTO	4 00	4.05	11- 444407	subunit
	GAATCCAACTG	1,00			similar to mouse neuronal protein 15.6
	CAGCGCGCCCT	1,00		Hs.152932	
1406	GCATAGGCTGC	1,00	4,69	Hs.12084	Tu translation elongation factor, mitochondrial
7407	ATCCCTCAGTG	1,00	1 06	He 1812/2	activating transcription factor 4 (tax-
' 40 /	AICCCICAGIG	1,00	4,50	1 13. 10 1243	responsive en
7408	CTGTTGGCATT	1,00	5.14	Hs 184108	ribosomal protein L21 (gene or
.50		.,50	٥, ١٦		pseudogene)
7409	CAGGAACGGGG	1,00	5,52	Hs.72241	mitogen-activated protein kinase kinase
					2
7410	GAAAAATTTAA	1,00	6,07	Hs.17775	p75NTR-associated cell death executor;
					ovarian granu
$\overline{}$	GACTCTTCAGT	1,00			alpha-1-antichymotrypsin
	CCTGGTCCCAA	1,00			keratin 7
7413	GGGGACTGAAG	1,00	6,93	Hs.3709	low molecular mass ubiquinone-binding
744	00000070700	4.00	0.00	11 0070	protein (9.5kD
<b> </b> /414	GGGCGCTGTGG	1,00	8,83	Hs.8372	ubiquinol-cytochrome c reductase
7/15	TCACCCAATAA	2.00	17.00	Un 02040	(6.4kD) subunit
	TGAGGGAATAA	2,00			triosephosphate isomerase 1
1410	GGAACAAACAG	1,00	9,05	ms.2/806/	Homo sapiens cDNA FLJ20161 fis, clone
7417	AACGACCTCGT	1,00	11 99	He 170661	COL09252, high tubulin, beta polypeptide
	AAGGGAGCACC	1,00			immunoglobulin lambda locus
	GCCGGGTGGGC	2,00			basigin
	CCTCCAGCTAC	2,00		Hs.242463	<del></del>
	CAAACCATCCA	1,00		Hs.65114	
11741		1,001	44.43	11 13.00 1 14	INGIAUII IO

Tabelle 5:

Nr.	Tag_Sequence	CGAP	Rel.	Quotient	signific.	UniGene	Beschreibung
			Expr. Haut			AccNr.	
1	ATCCGCGAGGC	0,12		375,00	63,99	Hs.180142	CLSP Calmodulin-like skin protein
2	GAGATAAATGA	0,09			31,88	Hs.3185	lymphocyte antigen 6 complex, locus D
	TAAACCTGCTG	0,47	110,00	234,04		Hs.99923	lectin, galactoside- binding, soluble, 7 (galectin 7)
- 4	GATGTGCACGA	1,08	216,00	200,00	289,47	Hs.117729	keratin 14 (epidermolysis bullosa simplex, Dowling-M
5	ACATTTCAAAG	0,00	161,00	161,00	248,57	Hs.80828	keratin 1 (epidermolytic hyperkeratosis)
6	TTTGTAGAGGA	0,26	37,00	142,31	47,97	Hs.279671	katanin p60 (ATPase- containing) subunit A 1
7	ACCTCCACTGG	0,00	139,00	139,00	214,56	Hs.112457	ESTs
8	AATCTTGTTTC	0,88	93,00	105,68	115,24	Hs.32343	ESTs
	GAAAACAAAGT	4,48	467,00	104,24	100,00	Hs.99936	keratin 10 (epidermolytic hyperkeratosis; keratosis
	GCCCCTGCTGA	1,46	148,00	101,37	181,69	Hs.195850	keratin 5 (epidermolysis bullosa simplex, Dowling-Me
11	CACACGGGCGA	0,26	26,00	100,00	32,16	Hs.194679	WNT1 inducible signaling pathway protein 2

Tabelle 4:

Nr.	Tag_Sequence	CGAP	Expr.	Quotie nt	signific.	UniGene AccNr.	Beschreibung
			Haut				
12	GACAATAAATG	0,15	9	60,00	10,38	Hs.137556	Homo sapiens mRNA; cDNA DKFZp434A132 (from clone DKF
13	ACTACCATAAC	0,12	7	58,33	8,06	Hs.57929	slit (Drosophila) homolog 3
14	CATTGTAAATA	0,12	. 7	58,33		Hs.55279	protease inhibitor 5 (maspin)
15	ACCGGCGCCCG	0,5	29	58,00			tetranectin (plasminogen-binding protein)
16	CCACCACGCTT	0,29	16	55,17	17,84		ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY
17	CCCCGGCCACC	0,88	42	47,73	44,79		(Manual assignment) desmin, muscle intermediate fila
. 18	TGAAATAAAAG	0,26	12	46,15	12,9	Hs.48516	ESTs
19	ACTGAGTAGGT	0,09	4	44,44	·		ATP-binding cassette, sub-family A (ABC1), member 8
20	ATCCTTGCTGA	0,7	. 26	37,14	26,12	Hs.2621	cystatin A (stefin A)
21	CAGCCTGGGTG	0,15	- 5	33,33	5,1	Hs.171941	
	GATATGTTATA	0,15	5	33,33			collagen, type XVII, alpha 1
	TGGCTTCATCA	0,12	4	33,33		Hs.646	carboxypeptidase A3 (mast cell)
	CCTGTAACACC	0,09		33,33			periplakin
	CCCCGGAGGTC	0,09		33,33			coagulation factor X
	AGATCAGTTGA	0,09	3			Hs.191805	
	CCCTCAGCACC	0,29		31,03			annexin A8
	CTTTATTCCAG	1,7		28,82			collagen, type I, alpha 1
	TCCACTGGCCT	0,82	23	28,05	21,32		ESTs
	CACGCAGTGGC	0,18	5	27,78 25,00	4,85	Hs.245545	
	TACATTATATA	0,12				Hs.198862	
	ATGGATACGGG	0,41		24,39			(Manual assignment) unclear, probably reverse tag o
	CCGGGGGAGCC	1,93		22,28	36,36	Hs.172928	collagen, type I, alpha 1
	CAGTTTTTTC	0,09		22,22	1,84	Hs.99597	ESTs
	GTGGATTCAAG	0,09		22,22		Hs.93847	NADPH oxidase 4
	TGTCTGTGTGT	0,09		22,22			ESTs
37	TCTACACGTGC	0,09	2	22,22	1,84	Hs.53155	properdin P factor,

			T			I	complement
30	GAAATGCCACT	0.00	2	22.22	4 04	Un 20052	
	GAAATGCCAGT	0,09		22,22		Hs.30853	ESTs
39	ACGAAACCTCG	0,09	2	22,22	1,84	ms.285/85	Homo sapiens cDNA
	.·						FLJ20115 fis, clone COL05594
40	GGCAATGCAGT	0,09	2	22,22	1 0/	Hs.275505	
	CCTTTTCAGCA		2	22,22		Hs.25930	
		0,09	2				ESTs
	CCTCTTTAACA	0,09		22,22		Hs.25750	ESTs
	TATCTAGCTGC	0,09	2				hypothetical protein
44	GCTGTAATCCT	0,09	2	22,22	1,84	HS.241382	tumor necrosis factor
	,					•.	(ligand) superfamily,
45	0000400000	- 0 00		00.00	4.04	11- 00500	member 1
45	GGGCAGCCGCC	0,09	2	22,22	1,84	Hs.23598	CREB binding protein
			Ρ,				(Rubinstein-Taybi
46	CGCTTGTTTAA	0.00		22.22	1 04	Ua 190209	syndrome)
40	CGCTIGTTIAA	0,09	2	22,22	1,04	IUS. 100380	LIM domain-containing
						·	preferred translocation partne
47	GCACACACCTG	0,09	2	22.22	1 04	Un 171112	EST, Weakly similar to
47	GCACACACCIG	0,09	2	22,22	1,04	ПS. 17 1 143	ALU1_HUMAN ALU
		:	•				SUBFAMILY J SE
48	CTGGAGTCGGC	0,09	2	22,22	1.8/	He 166371	Interleukin-1
70	01004010000	0,03	-	22,22	1,04	118.100371	Superfamily z
49	GAGGTCAGTTG	0,09	2	22,22	1 84	Hs 151696	DKFZP727G051 protein
	CCAGGCAAGAC	0,09	2	22,22			distal-less homeo box 3
	GAAATCAAAAA	0,59	13	22,03			sialic acid binding Ig-like
L	5/00/10/000	0,00	13	22,00	11,5	118.117000	lectin 5
52	AATCTAGTTCT	0	22	22,00	33,72	Hs.251440	Human profilaggrin gene
							exons 1-3, 5' end
53	AAGCTAATAAA	0,41	9	21,95	7,89	Hs.88474	prostaglandin-
٠.		.					endoperoxide synthase
							1 (prostaglandin
54	TGTGCGGCTTC	0,23	5	21,74	4,43	Hs.162196	hypothetical protein
							FLJ20321
55	CAGGTTTCATA	3,08	66	21,43	54,72	Hs.24395	small inducible cytokine
						. ,	subfamily B (Cys-X-
<u> </u>	07070070470						Cys), me
56	CTGTCGTCATC	0,35	.7	20,00	5,98	Hs.183860	hypothetical protein
	47400460706						FLJ20277
57	ATAGCACGTGC	0,15	3	20,00	2,66	Hs.277329	ESTs, Weakly similar to
							ALU5_HUMAN ALU
	OTOAOA A OTOG	- 245		00.00		11 05000	SUBFAMILY SC
58	GTGAGAACTCG	0,15	3	20,00	2,66	Hs.250639	ES   S

Tabelle 3:

Nr.	Tag_Sequence	CGAP	Rel.	Quotient	signific.	UniGene	Beschreibung
			Expr.		0.90.	AccNr.	1200,0
			Haut				
59	ACTTATTATGC	1,49	29	19,46	23,46	Hs.76152	decorin
60	CTTGCAGTCCT	0,26	5	19,23		Hs.27018	Ris
61	CATCTGTACTC	0,73	14	19,18	11,49	Hs.180255	major histocompatibility
							complex, class II, DR
٠							beta
	GTGGAGGGCAC	1,03				Hs.83393	cystatin E/M
63	AGGCAGGAAAA	0,29	5	17,24	4,09	Hs.133081	ESTs, Weakly similar to
]							hypothetical protein
		0.50		10.00	= 0.1		[H.sapi
64	AATTGAAAAGG	0,59	10	16,95	7,94	Hs.78344	myosin, heavy
- 1							polypeptide 11, smooth
65	CTTTAAAATGA	0.10	3	16.67	2.5	Un 9247	muscle
	TGTGCCAGTTT	0,18 0,18	3	16,67 16,67			stromal antigen 2
00	TOTOCCAGTIT	0,10	3	10,07	2,5	ms.55556	ESTs, Weakly similar to ALU8 HUMAN ALU
							SUBFAMILY
67	AGTAGCTGGGA	0,18	3	16,67	2.5	He 22/53/	EST, Weakly similar to
۱'`	ACIACOICCA	0, 10		10,07	2,5	1 15.224554	alternatively spliced
				İ			product
68	CCTCTGTCTCC	0,18	3	16,67	2.5	Hs 161031	Homo sapiens mRNA;
		,		, , , , ,	_,_		cDNA DKFZp434K0322
							(from clone
69	AACATTTAGGA	0,18	3	16,67	2,5	Hs.138380	KIAA0624 protein
70	CAATAAAATTT	0,18	3	16,67			tumor protein 63 kDa
					-		with strong homology to
	<u></u>						p53
	GCCGCTCAAGG	0,18	3	16,67		Hs.126064	
72	CCTGGTCAAGA	0,12	2	16,67	1,68		silver (mouse homolog)
							like
73	CCACCGCAGGA	0,12	2	16,67	1,68	Hs.85112	insulin-like growth factor
74	ACACTTOTOAA	0.40		40.07	4.00		1 (somatomedia C)
/4	ACACTTCTCAA	0,12	2	16,67	1,68		glutathione S-
75	CCTCTCTGGTC	0.12	2	16.67	1.60	Un 50074	transferase M5
13	COTOTOTOGIC	0,12	2	16,67	1,00		heat shock 27kD protein
							family, member 7 (cardiovasc
76	GCATATCTGTG	0,12	2	16,67	1 68	Hs.5459	KIAA1436 protein
	AGCTGTGATGG	0,12	2	16,67		Hs.249983	
	GCTAACTTAAA	0,12	2	16,67			ESTs
	CCTTGAAATCA	0,12	2	16,67		Hs.183161	
	CTITATCAATA	0,12	2	16,67			microphthalmia-
	- : · · · · · · · · · · · · · · · · · ·	٥, ٠٠-		.5,57	1,00	100017	associated transcription
							factor
81	ACAGCCCTGAT	0,12	2	16,67	1.68		ribosomal protein L18a

82	GATACTCAGAA	0,12	2	16,67	1.68	Hs.144726	ESTs
	GCCTGGGAGAC	0,12	2	16,67		Hs.118346	
	TGGGTGGTGGT	0,79	13	16,46			fragile X mental
					·		retardation, autosomal
							homolog 1
85	AGCTACCACAG	0,59	9	15,25	6,87	Hs.169886	tenascin XB
86	ACAGCGGCAAT	4,63	69	14,90	49,04	Hs.74316	desmoplakin (DPI, DPII)
87	GTAAAATCCCA	0,21	3	14,29	2,36	Hs.278623	ESTs, Weakly similar to
Ì			İ				ALU1_HUMAN ALU
							SUBFAMILY
	CACTTGTAATC	0,21	3	14,29			KIAA1185 protein
	CTTGTAGTTCC	0,21	3	14,29			KIAA0677 gene product
90	GGGTTTTCTGG	0,21	3	14,29	2,36	Hs.153703	ESTs, Moderately
		1	[			!	similar to
			l				DHSA_HUMAN
							SUCCINATE
	CAGCAGAACTG	0,21	3	14,29			CGI-43 protein
92	CCACAGGAGAA	5,68	81	14,26	56,23	Hs.169902	solute carrier family 2
		ľ					(facilitated glucose
	ATA 000 A 000 A	0.00		10.70			transpo
93	ATAGCCAGGGA	0,29	4	13,79	3,02	Hs.95582	SRY (sex determining
	OTA CA A A A OTA			40.70	0.00	11. 0550	region Y)-box 20
	GTACAAAAGTA	0,29	4	13,79			binder of Arl Two
	TCACAGGGTCC	0,29	4	13,79			lamin A/C
_	TTCTGTGTGCC	0,29	4	13,79			ESTs
	TAGCCGGGACG	0,73	10	13,70			Kruppel-like factor 2 (lung)
98	ATCACACAGCT	0,44	6	13,64	4,44	Hs.79386	leiomodin 1 (smooth
							muscle)
99	ATCTCGAAAGG	0,59	8	13,56	5,84	Hs.10784	hypothetical protein
<u> </u>					····		FLJ20037
100	GACCCAACTGG	0,15	2	13,33	1,54	Hs.89575	CD79B antigen
				-			(immunoglobulin-
404	04040004004	0.45		10.00	4.54	11 0 1770	associated beta)
	CACAGGGAGGA	0,15	2	13,33			KIAA0246 protein
	CAGCTGGCCCA	0,15	2	13,33			fibulin 1
103	GAGGGCTTTGC	0,15	2	13,33	1,54		aldo-keto reductase
							family 1, member C3 (3-
104	GTGAGCCAACA	0.45		12 22	4 5 4		alpha hyd
104	GTGAGCCAAGA	0,15	2	13,33	1,54	Hs.75410	heat shock 70kD protein
				j			5 (glucose-regulated protein
105	TACCCCAAAAA	0,15	2	13,33	1 54	Hs.6449	hypothetical protein
							FLJ20542
	CAGGATGCTTG	0,15	2	13,33	1,54	Hs.56729	lymphocyte-specific protein 1
107	AGTGTGTTGCA	0,15	2	13,33	1,54	Hs.56105	ESTs, Weakly similar to
			- 1				WDNM_RAT WDNM1
1.55	400400=======				<del></del>		PROTEIN
108	AGGACCTGAAG	0,15	2	13,33	1,54	Hs.32352	hypothetical protein

						<del></del>	DVE7-424V1240
400	ATTOTOGTOAT	0.45		40.00	4.54	11- 40070	DKFZp434K1210
109	ATTCTGGTCAT	0,15	2	13,33	1,54	Hs.18878	ESTs, Weakly similar to
113	000000000000000000000000000000000000000	0.45		- 40.00		11 100000	dJ876B10.4 [H.sapiens]
	CCTTTTGGGAG	0,15	2	13,33		Hs.186600	
111	CGGTTCATACA	0,15	2	13,33	1,54	Hs.169487	Kreisler (mouse) maf-
			j				related leucine zipper
							homolog
112	ATGGTGCCACC	0,15	2	13,33	1,54	Hs.161554	hypothetical protein
							FLJ20159
113	TAATGTTAATG	0,15	2	13,33	1,54	Hs.153924	death-associated ·
							protein kinase 1
114	AGGGCCCTCTG	0,15	2	13,33	1,54	Hs.129014	hypothetical protein
							FLJ20207
115	GTGGCACGCGT	0,15	2	13,33	1,54	Hs.118243	deoxyribonuclease II,
							lysosomal
116	AGCTTGAGTTC	0,15	2	13,33	1,54	Hs.117582	CGI-43 protein
	GTGGGGCCAAG	0,53	7	13,21			folate receptor 2 (fetal)
	GTGAAGCCTCA	0,38	5	13,16		Hs.271823	
	ACCAGACAGAC	0,23	3	13,04			ESTs
	GTGAAACTCTT	0,23	3	13,04			RNA POLYMERASE I
'2	010/00/01011	0,20	ĭ	10,04	2,20	110.20700	AND TRANSCRIPT
			.				RELEASE
121	ATTTCCATTAA	0,23	3	13,04	2 23	He 28/126	hairless (mouse)
'2'	ATTIOCATION	0,23	ી	13,04	2,25	115.204120	homolog
122	GTGGTAAGCAC	0,23	3	13,04	2 22	Un 271927	ESTs, Moderately
122	GIGGIAAGCAC	0,23	ી	13,04	2,23	ПS.21 1021	similar to
							ALU7_HUMAN ALU
122	GTTTTGCCCAC	0,23	3	13,04	2 22	Ho 151407	cartilage intermediate
143	GITTIGCCCAC	0,23	ી	13,04	2,23	ПS. 15 1407	
							layer protein, nucleotide
124	GCCCACACAGC	0.62	0	42.00	E 74	Un 1600	pyr
124	GCCCACACAGC	0,62	8	12,90	5,71	Hs.1690	heparin-binding growth
125	TTTCCTCTCAA	2.00	20	40.74	25.2	11- 404540	factor binding protein
	TTTCCTCTCAA	2,99	38	12,71		Hs.184510	
126	CGGGAGCGCTA	1,05	13	12,38	8,84	HS.148590	ESTs, Weakly similar to
			Ī				AF208846_1 BM-004
407	TTOOATATOAG	0.00	40	40.00	0.04	11 00007	[H.sapiens
127	TTGCATATCAG	0,82	10	12,20	6,84	Hs.82237	ataxia-telangiectasia
	•	ŀ		ļ			group D-associated
405	A0000T0000			45.5		11 00000	protein
128	AGGCCTCGGCA	0,41	5	12,20	3,55	Hs.286202	Homo sapiens cDNA
							FLJ11346 fis, clone
						<u> </u>	PLACE1010900
	GTGGCGAATGA	0	12	12,00		Hs.69752	desmocollin 1
130	TGTGAAGCCTT	0	12	12,00	18,26	Hs.5476	serine protease
							inhibitor, Kazal type, 5
131	TCAGACTTTTG	0,76	9	11,84	6,1	Hs.5889	ESTs, Weakly similar to
							AC004876_5 similar to
							predic
132	ATTTCTTCAAG	0,76	9	11,84	6,1	Hs.31386	ESTs, Highly similar to
						ĺ	JE0174 frizzled protein-
	•		•			•	•

<u></u>							2 -
133	GAATTATACTT	0,85					hypothetical protein FLJ10134
134	TCTGGGGAACA	0,26	3	11,54	2,12	Hs.184390	similar to aspartate beta hydroxylase (ASPH)
135	GCAAAAACCCG	0,26	3	11,54	2,12	Hs.184109	ribosomal protein L37a
136	AATGTTGTGCA	0,35	4	11,43			cytochrome P450
				·			retinoid metabolizing protein
137	ACAATGTTGTA	0,18	2	11,11	1,43	Hs.7678	cellular retinoic acid- binding protein 1
138	CGAGAGTGTGA	0,18	2	11,11	1,43	Hs.58210	ESTs
139	GTATAAAAAAA	0,18	2	11,11	1,43	Hs.27337	hypothetical protein FLJ20623
140	AGGTCGAGGCT	0,18	2	11,11	1,43	Hs.270125	
	CCCGGCCCAGT	0,18	2	11,11			EST, Weakly similar to ALUA_HUMAN !!!! ALU CLASS A
142	TTGACCCAGCC	0,18	2	11,11	1,43	Hs.193745	ESTs
143	TATTTTATTTG	0,18	2	11,11			purinergic receptor (family A group 5)
144	GCATCATAGGT	0,18	2	11,11	1,43	Hs.184108	ribosomal protein L21 (gene or pseudogene)
145	TACCGCTCCCT	0,18	2	11,11	1,43	Hs.172803	Homo sapiens mRNA; cDNA DKFZp434G2416 (from clone
146	СТССТӨТӨӨТС	0,18	2	11,11	1,43	Hs.169851	ESTs, Weakly similar to ALU4_HUMAN ALU SUBFAMILY
147	GGTGTCTCCTC	0,18	. 2	11,11	1,43	Hs.146038	
148	CAATCTTGTGA	0,18	2	11,11		Hs.104353	
149	CCTACAAAAA	0,09	1	11,11			ESTs
150	TGTGCCCAGCC	0,09	1	11,11		Hs.97905	ovo (Drosophila) homolog-like 1
151	GATGGGGACAG	0,09	1	11,11	0,69	Hs.92195	ESTs
152	TAAAAATATTG	0,09	1	11,11			insulin receptor
153	TTTAAGTTAGG	0,09	1	11,11		Hs.82932	cyclin D1 (PRAD1: parathyroid adenomatosis 1)
	GTATATGTATT	0,09	1	. 11,11	0,69	Hs.7917	DKFZP564K247 protein
	CTAAAGTGTCA	0,09	1	11,11		Hs.7910	RING1 and YY1 binding protein
	AATTTGGCTTT	0,09	. 1	11,11	0,69		Homo sapiens cDNA FLJ20684 fis, clone KAIA3469
	CCGTGGCACCA	0,09	1	11,11			ESTs
	AGCACCCTTGT	0,09	1	11,11			protein kinase C binding protein 1
	CCTTTGAGAGC	0,09	1	11,11	0,69	Hs.71791	hypothetical protein
160	TAAATGTAAAT	0,09	1	11,11			KIAA0427 gene product

161	TACAGACATAC	0,09	1	11,11	0,69	Hs.63984	cadherin 13, H-cadherin (heart)
162	GTAGCATTTGC	0,09	1	11,11	0,69	Hs.63302	myotubularin related protein 3
	AAGTAGGTTTT	0,09	1	11,11	·	Hs.50216	zinc finger protein (ZFD25)
164	ATCACTCCCCA	0,09	1	11,11	0,69	Hs.37058	calcitonin/calcitonin- related polypeptide, alpha
165	TAGGCAGACCT	0,09	1	11,11	0,69	Hs.35488	ESTs, Moderately similar to ALU6_HUMAN ALU
166	GCCTGGCCAGG	0,09	1	11,11	0,69	Hs.3343	phosphoglycerate dehydrogenase
	GTCATCTTGTT	0,09	1	11,11		Hs.32366	ESTs, Moderately similar to TWST_HUMAN TWIST
	CTCAACAACCA	0,09	1	11,11	·	Hs.30036	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY
169	TTATATTAATA	0,09	1	11,11	0,69	Hs.29205	alpha integrin binding protein 63
170	GATTAAACCTT	0,09	1	11,11			ESTs, Moderately similar to meningioma-expressed ant
171	GATGGAGGTTA	0,09	1	11,11	0,69	Hs.285224	ESTs, Weakly similar to unnamed protein product [H.s
172	TACAGGCGTGG	0,09	1	11,11	0,69	Hs.283329	
173	TGTGTGTGTAT	0,09	1	11,11			Homo sapiens mRNA; cDNA DKFZp434J1630 (from clone
	GGTGCCTGTAA	0,09	1	11,11			ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS C
	AAGTTGTGGCC	0,09	1	11,11			tubulin, alpha, ubiquitous
	ACAGGAGCGTT	0,09	1	11,11		Hs.275896	
	ATGCCTACTCT	0,09	1	11,11		Hs.274834	
	AGCACAATCTT	0,09	1	11,11			Homo sapiens cDNA FLJ10131 fis, clone HEMBA1003041
$\overline{}$	GTTCTGTTTGG	0,09	1	11,11		Hs.271040	
	TTGTGATTATA	0,09	1	11,11			TRABID protein
	TATGCAGATCA	0,09		11,11		Hs.254948	
	GTTGAGGACAT	0,09		11,11		Hs.250520	
	GCCAGTGGCTG	0,09	1	11,11		Hs.249720	
	GCAGATCTTTC	0,09	1	11,11		Hs.248623	
	TTGAAACTTCT	0,09	1	11,11		Hs.242481	
1 100	CAAAAAGTTGA	0,09	1	11,11	0,69	HS.241425	DKFZP586O1422

			Т	T			protein
187	CTCCATTCTCA	0,09	1	11,11	0.69	Hs.23803	Homo sapiens mRNA;
		-,55	1	,	3,00		cDNA DKFZp434M2217
			l		-		(from clone
188	GCCACAGCTGG	0,09	1	11,11	0.69	Hs.23565	ESTs
$\overline{}$	GGGAAAGAAGG	0,09	1	11,11		Hs.233193	
	CAAAAGAATAA	0,09	1	11,11		Hs.233013	
	TAACAGTAATA	0,09	1	11,11		Hs.231913	
	TGCCTAGGAAA	0,09	1	11,11			Homo sapiens mRNA;
'02		. 5,55	']	11,11	3,03	5.220000	cDNA DKFZp586F1922
							(from clone
193	CCCTAGGAGAC	0,09	1	11,11	0.69	Hs.217484	
	GGGTTGTTGTA	0,09	1	11,11		Hs.211258	
	AACCCCCAAAC	0,09	1	11,11		Hs.207181	
$\overline{}$	CTTTTTCTTT	0,09	1	11,11		Hs.204917	
	TGGGAATTGTG	0,09	1	11,11		Hs.203750	
	CATTCCAGCCT	0,09	1	11,11		Hs.201306	
	GCTGGAGTGCA		<del>-   </del>	11,11			Homo sapiens mRNA
199	GUIGGAGIGCA	0,09	'	11,11	0,09		full length insert cDNA
							clone EURO
200	GAAGGCCAGCT	0,09	1	11,11	0.60	Hs.194624	
	ACGTATTTGAG	0,09	1	11,11			Homo sapiens mRNA;
201	ACCIAITIGAG	0,09	·	11,11	0,09	113.133400	cDNA DKFZp434O1521
					ļ		(from clone
202	TAATTTAAACC	0,09	1	11,11	0.60		hypothetical protein
202		0,09	. '1	11,11	0,09		FLJ20281
203	CTAAATGTGAA	0,09	1	11,11	0.60		high-mobility group
200	01/0010/07	5,09	'	11,11	0,09		(nonhistone
							chromosomal) protein
204	CCTGGCTCTAA	0,09	1	11,11	0.69		Homo sapiens cDNA
- "		5,00	'	,.	3,00		FLJ11333 fis, clone
				1			PLACE1010616
205	CTGTCCTTGTT	0,09	1	11,11	0.69	Hs.176333	
	CAATATTTGAG	0,09	1	11,11			a disintegrin and
		-,	']		5,55		metalloproteinase
							domain 28
207	CAATGGATGGC	0,09	1	11,11	0.69		ESTs, Weakly similar to
		-,	1	,			AF126780_1 retinal
			ļ	ļ			short-cha
208	GTCCTTGACCA	0,09	1	11,11	0.69	Hs.170524	
	GCAATGACCTG	0,09	1	11,11			interleukin 9 receptor
	TCCTAATTCAG	0,09	1	11,11			KIAA1357 protein
	TTAATGATCTT	0,09	1	11,11		Hs.165240	
_	TTAAGTGTTCT	0,09	1	11,11			toll-like receptor 4
	TGAAGCGTTTA	0,09	1	11,11			protein tyrosine
- :	. =	5,00	']	,	3,55		phosphatase, non-
		ì		İ			receptor type 21
214	GTGGAGCTTAA	0,09	1	11,11	0.69	Hs.152385	
	GAGACTGGGGC	0,09	1	11,11		Hs.142854	
	GTGGTACTCGC	0,09	1	11,11			ESTs, Weakly similar to
1 - 10		0,00	4	1 1, 1 1	0,00	1.13.171040	LC 10, VVCakiy Similar to

							050504 :
1							S59501 interferon
047	TOOLOTOOTTO	0.00		44.44	0.00	11. 444660	receptor J
	TGCAGTGCTTG	0,09	1	11,11			chloride channel 2
$\overline{}$	AGCTTATTGGC	0,09	1	11,11			KIAA1098 protein
$\overline{}$	ACTTCTGCTTA	0,09	1	11,11		Hs.13740	
	GATAGAAATTT	0,09	1	11,11		Hs.131987	
	TTTGACTAATT	0,09	1	11,11		Hs.131761	
	TCAAAATGACA	0,09	1	11,11		Hs.131272	
	CCAGCTAGTTT	0,09	1	11,11		Hs.128692	
	TAGAGGAGTTG	0,09	1	11,11		Hs.125815	
-	GGGAAACACCA	0,09	1	11,11		Hs.123471	
226	GAAATGAGTGT	0,09	1	11,11	0,69	Hs.1200	arachidonate 12-
							lipoxygenase
	ATCTTGGTACT	0,09	1	11,11			fibronectin 1
	CATATCATCTC	0,09	1	11,11		Hs.118130	
	ATGTGACTTTT	0,09	1	11,11			CGI-43 protein
	CCACTGTAAGC	0,09	1	11,11			CGI-43 protein
231	TATAAGGCTGA	0,09	1	11,11	0,69		ESTs, Weakly similar to ZN84_HUMAN ZINC FINGER
232	GCCACTGCCAC	0,09	1	11,11	0,69	Hs.112405	S100 calcium-binding protein A9 (calgranulin B)
233	TGAGGCCAGGG	0,09	1	11,11	0,69	Hs.110128	hypothetical protein FLJ10060
234	GTTTATTTGAA	0,09	1	11,11	0,69	Hs.109087	ESTs
	AAAATTGTTAG	0,09	1	11,11	0,69	Hs.10760	hypothetical protein FLJ20129
236	GTGATGGGCTC	0,82	9	10,98	5,88	Hs.25482	envoplakin
237	AGGCTCCTGGC	4,81	52	10,81	31,57	Hs.24395	small inducible cytokine subfamily B (Cys-X- Cys), me
238	AACAGCAAGGA	0,47	5	10,64	3,33	Hs.20665	ESTs
	CCACGGGATTC	1,32	14	10,61	8,79	Hs.119571	collagen, type III, alpha 1 (Ehlers-Danlos syndrome
240	GATTTCGTTTT	0,38	4	10,53	2,68	Hs.738	early growth response 1
	TATAGCCCTCA	0,38	4	10,53			a disintegrin and metalloproteinase
242	CTCACACCTCC	0.30		10.50	0.00	U= 202000	domain 17 (tumor
	GTGACACGTGC	0,38	4	10,53		Hs.282996	
	AGCTGTCGTAG	0,29	3	10,34			ESTs Madamatah
244	ATTGTTTCAAG	0,29	3	10,34	2,02	Hs.32366	ESTs, Moderately similar to TWST HUMAN TWIST
245	AGACCCTGTCT	0,29	3	10,34	2 02	Hs.239283	
_	TTGGCAAGGCT	0,29	3	10,34		Hs.184720	
	ATCATAGCTCA	0,59	6				ESTs, Moderately
		0,03		- 10,17	0,07		similar to ALU1_HUMAN ALU

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Tabelle 2:

Nr.	Tag_Sequence	CGAP	Rel. Expr. Haut	Quotie nt	signifi c.	UniGene AccNr.	Beschreibung
248	CCTACCACCAT	1,11		9,91	6.73	Hs.8468	RelA-associated inhibitor
	CCAGGGCAACA	4,1				Hs.120980	(Manual assignment) ORF-less transcript in MEN1 regi
250	TACAGTATTTT	0,21	2	9,52	1,33	Hs.82921	solute carrier family 35 (CMP-sialic acid transporte
251	TGCCAGGTGCA	0,21	2	9,52	1,33	Hs.75442	albumin
	TTTTTATTCC	0,21				Hs.327	interleukin 10 receptor, alpha
253	AATATTTTAT	0,21	2	9,52	1,33	Hs.31386	ESTs, Highly similar to JE0174 frizzled protein-2
254	GAGAACCACCT	0,21	2	9,52	1,33	Hs.30250	v-maf musculoaponeurotic fibrosarcoma (avian) oncoge
255	GACACACCGAA	0,21	2	9,52	1,33	Hs.274243	receptor tyrosine kinase- like orphan receptor 1
256	GGCTTGTCTAT	0,21	2	9,52	1,33	Hs.23294	ESTs, Weakly similar to weak similarity to HSP90 [C.
257	GGCAATATAGT	0,21	2	9,52	1,33	Hs.194429	ESTs, Weakly similar to unknown protein [H.sapiens]
258	AGGATAACTTC	0,21	. 2	9,52	1,33	Hs.184482	DKFZP586D0624 protein
259	GATCAATCAGT	0,21	2	9,52	1,33	Hs.16530	small inducible cytokine subfamily A (Cys-Cys), memb
260	TTGAATATTAA	0,21	2	9,52	1,33	Hs.161554	hypothetical protein FLJ20159
	TCCTCACTTCA	0,21	2	9,52		Hs.158455	ESTs
	GTTGCAGCATT	0,21	2	9,52	1,33	Hs.147189	HYA22 protein
	GTGTCTGTCTC	0,21				Hs.137432	ESTs
_	GGCCTCTCCGA	0,21				Hs.132834	hematopoietic protein 1
	ACAGAATGCCT	1,79				Hs.79732	fibulin 1
**	GGGGCTGCCCA	1,58	15	9,49	8,82	Hs.195727	tumor endothelial marker 1 precursor
267	AAAATCGCTTG	0,53		9,43	3,14	Hs.226581	COX15 (yeast) homolog, cytochrome c oxidase assembly
268	GTATAAACGTC	0,32	3	9,38	1,93	Hs.237356	stromal cell-derived factor 1

							<del>, </del>
269	CACTITACCAG	0,32	3	9,38	1,93	Hs.170019	runt-related transcription factor 3
270	TTAATTACAGT	0,32	3	9,38	1,93	Hs.159640	serum/glucocorticoid regulated kinase
271	TCAGCGACCCT	0,44	4	9,09	2,49	Hs.169946	GATA-binding protein 3
	ATGGCACATTC	0,44	4	9,09	2,49	Hs.14328	Homo sapiens mRNA; cDNA DKFZp762O124 (from clone
	AGGCTCAGGTC	0	9	9,00		Hs.78344	myosin, heavy polypeptide 11, smooth muscle
274	AACAGGGGCCA	0,56	5	8,93	3,05	Hs.262958	ESTs, Weakly similar to alternatively spliced produc
275	CTGAAATCTAT	0,56	5	8,93		Hs.253467	ESTs
	GAGAAATCCCG	0,56	5	8,93		Hs.150298	ESTs
	TCAAAAGACCT	1,38	12	8,70		Hs.25647	v-fos FBJ murine osteosarcoma viral oncogene homolog
$\overline{}$	GAAAACAAACA	0,23	2	8,70		Hs.83004	interleukin 14
279	TGCATCTGTAC	0,23	2	8,70		Hs.58589	glycogenin 2
280	ATGAAACTCCA	0,23	2	8,70		Hs.277951	EST
281	TTTATTCCTCT	0,23	2	8,70	1,25	Hs.26290	ESTs
282	GGGAGACCTGT	0,23	2	8,70	1,25	Hs.256115	EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J
	CACTATGTAAA	0,23	2	8,70	·	Hs.24143	Wiskott-Aldrich syndrome protein interacting protein
284	TACAGCGGCAG	0,23	2	8,70		Hs.205616	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY
	AGTGCCTTGGG	0,23	2	8,70		Hs.178604	ESTs
	CGATGCTGACG	0,23	2	8,70		Hs.161554	hypothetical protein FLJ20159
	GTGGTGTAATC	0,35	3	8,57		Hs.261734	ESTs, Moderately similar to ALU7_HUMAN ALU
	TATCCCAGAAT	0,35	3	8,57		Hs.175819	EST
	TCTAAAAAGGC	0,35	3	8,57		Hs.16622	zinc finger protein 185 (LIM domain)
	TCGAAACGCTG	0,35	3	8,57	1,84	Hs.136528	ESTs, Moderately similar to ALU1_HUMAN ALU
	ACATTCTTTTT	2,58	22	8,53	11,94	Hs.82226	glycoprotein (transmembrane) nmb
292	CTGTTTGTTCA	0,94	8	8,51	4,59	Hs.211582	myosin, light polypeptide kinase
293	CAGTACTGTAT	0,47	4	8,51	2,41	Hs.9295	elastin (supravalvular aortic stenosis, Williams- Beu
294	GGTGAAACCCC	0,47	4	8,51	2,41	Hs.284878	EST

000	00044400004	0.47		0.54	0.44	11- 404004	IFOT-
	GCCAAACCCCA	0,47	4	8,51		Hs.194264	ESTs
	GCCGACGCCAG	0,47	4	8,51		Hs.165565	ESTs
	GAAGAGGACAA	0,59	5	8,47		Hs.120451	ESTs
298	TAAATAAGAAA	0,12	1	8,33	0,61	Hs.99875	zona pellucida binding protein
299	AAAGATCCCTC	0,12	1	8,33	0,61	Hs.94998	ESTs
300	ATCAAAGGTTA	0,12	1	8,33	0,61	Hs.94795	Homo sapiens mRNA; cDNA DKFZp564O222 (from clone
301	GCAAGGTTGGT	0,12	1	8,33	0,61	Hs.94761	ESTs, Weakly similar to KIAA0561 protein [H.sapiens]
302	GGGACAAAAA	0,12	1	8,33	0,61	Hs.93788	ESTs
303	ACTACCTCTGA	0,12	1	8,33	0,61	Hs.92254	hypothetical protein FLJ20163
304	GCTGGGCCCAG	0,12	1	8,33	0,61	Hs.90964	Homo sapiens cDNA FLJ20812 fis, clone ADSE01316
305	CAAAGAAATAG	0,12	1	8,33	0,61	Hs.88653	ESTs
306	AATACAAGTAT.	0,12	11	8,33		Hs.8707	KIAA1301 protein
307	CTCTGTGGCTC	0,12	1	8,33	0,61	Hs.85112	insulin-like growth factor 1 (somatomedia C)
308	TATTTGAAAGT	0,12	1	8,33	0,61	Hs.82664	ETAA16 protein
309	CCCGCCTCCGT	0,12	1	8,33	0,61	Hs.82071	Cbp/p300-interacting transactivator, with Glu/Asp-ri
310	TCACCGTAGEC	0,12	1	8,33	0,61	Hs.82042	solute carrier family 23 (nucleobase transporters),
311	CCAAAAATTAA	0,12	1	8,33	0,61	Hs.81424	ubiquitin-like 1 (sentrin)
312	ACTCGTATATG	0,12	1	8,33	0,61	Hs.81134	interleukin 1 receptor antagonist
313	TCAAGATGAAG	0,12	1	8,33	0,61	Hs.78948	Rab geranylgeranyltransferas e, beta subunit
	CGTCTATCCAT	0,12	1	8,33		Hs.76084	lamin B2
315	AAAGTGAAATG	0,12	1	8,33	0,61	Hs.75912	KIAA0257 protein
	TTCTGGTGCTG	0,12	1	8,33		Hs.75725	transgelin 2
317	CTGGCGTCGTC	0,12	1	8,33	0,61	Hs.75640	natriuretic peptide precursor A
	GGCTGGGGAGG	0,12	1	8,33	0,61	Hs.75061	MARCKS-like protein
319	TGCCCCAAAA	0,12	1	8,33	0,61	Hs.74649	cytochrome c oxidase subunit VIc
320	TTTATTGAACA	0,12	1	8,33	,	Hs.74649	cytochrome c oxidase subunit VIc
321	CTTTCCTCATT	0,12	1	8,33	0,61	Hs.7381	voltage-dependent anion channel 3
322	TTGGACTGAGG	0,12	1	8,33	0,61	Hs.6518	ganglioside expression factor 2
323	AGCATTAAAAA	0,12	1	8,33	0,61	Hs.61638	myosin X



324	CTGCTTTAAAA	0,12	4	8,33	0.61	Hs.56023	brain-derived
324	CIGCILIAAAA	0, 12	'}	0,33	0,01	H8.56023	neurotrophic factor
325	TGCTTAAAAAT	0,12	1	8,33	0.61	Hs.5534	ESTs .
	ATGAACCCCCT	0,12	1	8,33		Hs.5011	RNA binding motif
020	/110/11000001	0,12	1	0,00	0,01	10.0011	protein 9
327	TGAAACTTCCC	0,12	1	8,33	0.61	Hs.4994	transducer of ERBB2, 2
	CAAACAAAAA	0,12	1	8,33		Hs.43728	hypothetical protein
	GAGGTAACTAC	0,12	1	8,33		Hs.43712	ESTs
	CATTAAAAAAAT	0,12	1	8,33		Hs.36908	activating transcription
	•						factor 1
331	CAATGGTGAAA	0,12	1	8,33	0,61	Hs.35093	lymphoid blast crisis
	A-1.1						oncogene
332	CTAATGAATGT	0,12	1	8,33	0,61	Hs.29809	Homo sapiens mRNA;
			İ				cDNA DKFZp434C185
222	ACCCCTTTCAC	0.40		0.33	0.64	He 20707	(from clone
	AGGGCTTTCAC TAACCGTGGAA	0,12	1 1	8,33 8,33		Hs.29797 Hs.29647	ribosomal protein L10 uncharacterized
334	IAACCGIGGAA	0,12	'}	0,33	0,01		hematopoietic
			i				stem/progenitor cells
335	TCTCCCACACC	0,12	1	8,33	0.61	Hs.2961	S100 calcium-binding
	101000/10/100	0, 12		0,00	. 0,0 !	110.200	protein A3
336	CCTGGATCTCC	0,12	1	8,33	0,61	Hs.28501	ESTs
	GGCCTGGCACT	0,12	1	8,33		Hs.283388	ESTs
	ACTITGTTTTT	0,12	1	8,33		Hs.28219	protein phosphatase 2
			j	[	•		(formerly 2A), regulatory
							subu
	GTGACGCCTGT	0,12	1	8,33		Hs.279361	EST
340	CCATAAGTCCT	0,12	1	8,33	0,61	Hs.279009	seven in absentia
-	1000000000	0.15					(Drosophila) homolog 1
341	AGCCCCCGCGC	0,12	1	8,33	0,61	Hs.272320	Homo sapiens mRNA;
		ŀ					cDNA DKFZp434L1226 (from clone
3/12	TATCATCATTC	0,12	1	8,33	0.61	Hs.270877	ESTs
	TTCCCTGAGCA	0,12	1	8,33		Hs.26198	ESTs, Weakly similar to
575	110001GAGUA	0, 12	'	0,00	0,01	10.20100	ALU1_HUMAN ALU
		[					SUBFAMILY
344	TAGTCTAAGGC	0,12	1	8,33	0,61	Hs.261782	ESTs
	ATAGTAGTAAT	0,12	1	8,33		Hs.258863	EST
	GATTTTAAATG	0,12	1	8,33		Hs.25333	interleukin 1 receptor,
							type II
	CCTGGCTAACA	0,12	1	8,33		Hs.252124	ESTs
	CTGTACAGACC	0,12	1	8,33		Hs.251653	tubulin, beta, 2
	GGCACCAGAGC	0,12	1	8,33		Hs.249614	EST
	GCGAACTCCGT	0,12	1	8,33		Hs.248844	ESTs
	CGCGTCCGTGT	0,12	1	8,33		Hs.243929	ESTs
352	GTTTTGGTTTA	0,12	1	8,33	0,61	Hs.241336	Homo sapiens mRNA;
						1	cDNA DKFZp564G0422
252	CTTTTTCCCAC	0.40		0.00	0.04	Un 040465	(from clone
	CTTTTTGCCAC	0,12	1	8,33		Hs.240165	ESTs Weekly eighter to
354	GTGGGGGGCGC	0,12	1	8,33	0,61	Hs.240031	EST, Weakly similar to

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			<del></del>				
					_		ALU2_HUMAN ALU SUBFAMILY S
355	TCTTTCCAACT	0,12	1	8,33	0,61	Hs.22394	hypothetical protein FLJ10893
356	TAGTAGGGCTC	0,12	1	8,33	0,61	Hs.21914	ESTs
357	AGGACATAACA	0,12	1	8,33	0,61	Hs.213793	ESTs
358	TTGTTTTAAGA	0,12	1	8,33		Hs.211519	hypothetical protein LOC56757
359	GTGGCACATCT	0,12	1	8,33		Hs.208925	ESTs, Weakly similar to alternatively spliced produc
360	CACATTGAGGC	0,12	1	8,33	0,61	Hs.207122	EST
	AGGCTAGCACT	0,12	1	8,33		Hs.206259	Homo sapiens mRNA for KIAA1190 protein, partial cds
	CCCCTGCCCTC	0,12	1	8,33		Hs.203317	EST
	TTGTCTCTTGA	0,12	1	8,33		Hs.20104	ESTs
	TGGGTACACTG	0,12	1	8,33		Hs.200030	ESTs, Weakly similar to ALUB_HUMAN !!!! ALU CLASS B
	GGCAGTGGTAA	0,12	1	8,33		Hs.197075	ESTs
	CATAAATGTTA	0,12	1	8,33		Hs.19479	ESTs
367	AGATTACCCAC	0,12	1	8,33		Hs.192155	ESTs
368	ACTGGCTCAGG	0,12	1	8,33	0,61	Hs.190719	ESTs
369	CTTGTAGTCTC	0,12	1	8,33	0,61	Hs.189073	ESTs
370	AGCCCAGCTGG	0,12	1	8,33	.0,61	Hs.18857	ESTs
371	GAGGGCAATCT	0,12	1	8,33	0,61	Hs.186753	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY
	CTGAAACAGGA	0,12	1	8,33	0,61	Hs.183601	regulator of G-protein signalling 16
	GCCCTACCTGC	0,12	1	8,33	0,61	Hs.182740	ribosomal protein S11
	GCCAACGGCGT	0,12	1	8,33		Hs.181002	MLL septin-like fusion (NOTE: non-standard symbol an
	GTAAAGATGAA	0,12	1	8,33		Hs.175941	B-cell receptor- associated protein BAP29
	GGGGTAATTTT	0,12	. 1	8,33	·	Hs.173497	Sec23 (S. cerevisiae) homolog B
	AGGACAATGAA	0,12	1	8,33	Ĺ	Hs.173135	dual-specificity tyrosine- (Y)-phosphorylation regula
	TTTGCACTTTT	0,12	1	8,33		Hs.167114	ESTs, Highly similar to AF070470_1 SPARC- related pro
	CTGAAGTGCAG	0,12	1	8,33		Hs.166609	ESTs
	CCCATTCAGTC	0,12	1	8,33		Hs.161554	hypothetical protein FLJ20159
381	TAATGTTCTCA	0,12	1	8,33	0,61	Hs.160271	G protein-coupled

	I	<del> </del>	— т	<del></del> -		T	recentor 49
202	CTTTOTCCACA	040	<del>-</del>	0.00	0.04	11- 450042	receptor 48
362	GTTTCTGCAGA	0,12	1	8,33	0,01	Hs.159642	glucosaminyl (N-acetyl) transferase 1, core 2 (beta-
383	GCAGATTCTCA	0,12	1	8,33	0.61	Hs.157716	ESTs
	GGAATGAGGGG	0,12	1	8,33		Hs.156452	ESTs
	GAGAGGAACTA	0,12	1	8,33		Hs.153523	ESTs
	TTCGACAGGCT	0,12	<del>-  </del>	8,33		Hs.152925	KIAA1268 protein
	CTCCAGCCTGG		<del> </del>			Hs.145331	
307	CICCAGCCIGG	0,12		8,33			ESTs, Weakly similar to ALUF_HUMAN !!!! ALU CLASS F
	AGTCTGTATTT	0,12	1	8,33		Hs.144906	ESTs
	GAGTATTATTT	0,12		8,33	0,61	Hs.143738	ESTs, Weakly similar to EPS8_HUMAN EPIDERMAL
390	TACTGTACTCC	0,12	1	8,33	0,61	Hs.143198	hypothetical protein similar to tumor
204	AAAATAAATO	0.40		0.00	0.04	11- 440000	suppressor p33
	AAAATAAAATG	0,12	1	8,33		Hs.142908	E2F-like protein
	CCCAGGAGTTT	0,12	1	. 8,33		Hs.13785	ESTs
	AAAATACAGTG	0,12	1	8,33		Hs.136433	ESTs
	CATTTGGCCGG	0,12	1	8,33		Hs.136031	ESTs
	CAGGACCTGAA	0,12	1	8,33		Hs.135971	ESTs
	TCATTAACAAA	0,12	1	8,33		Hs.135260	ESTs
	CAGTAGGATAA	0,12	1	8,33		Hs.134541	ESTs
	GCCTCACCTGG	0,12	1	8,33		Hs.128514	ESTs
- 4	TCTGTATCAAA	0,12	1	8,33		Hs.128408	ESTs, Moderately similar to ALU1_HUMAN ALU
	TGATTTGTGAA	0,12	1	8,33		Hs.12282	ESTs
	AGGATATTGGA	0,12	1	8,33		Hs.117721	ESTs
	GTGGTACACAG	0,12	1	8,33		Hs.117582	CGI-43 protein
	TTGGCCAGATT	0,12	1	8,33	0,61	Hs.117582	CGI-43 protein
404	ATTCTTGTACA	0,12	1	8,33	0,61	Hs.117527	ESTs
405	CCTTCTTGGGG	0,12	1)	8,33	0,61	Hs.117474	ESTs
406	TACTGGAAGGC	0,12	1	8,33		Hs.116874	ESTs, Weakly similar to putative p150
407	GTGGTGGTTGG	0,12		8,33	0.64	Hs.114408	[H.sapiens]
	AGTAATGAAAA		- 1				toll-like receptor 5
	TTAGTTTTGCT	0,12	1	8,33		Hs.11217	KIAA0877 protein
		·		8,33		Hs.108885	collagen, type VI, alpha
	GCTAATATATT	0,12	1	8,33		Hs.107883	ESTs
	TCTTGACTCCC	0,12	1	8,33	<u>-</u>	Hs.107265	ESTs
	TGTACTTAATT	0,12	1	8,33		Hs.107082	ESTs, Moderately similar to alternatively spliced pr
413	CTCAGCAGGAG	0,12	1	8,33	0,61	Hs.105489	ESTs, Weakly similar to AF109127_1 stromal cell-deri
414	GGTACAATCCG	0,12	1	8,33	0,61	Hs.104557	hypothetical protein

			7				FLJ10697
415	ACTCCAGACCT	0,12	1	8,33	0.61	Hs.104350	ESTs
	GAGAAACTCCG	1,08	9	8,33		Hs.184367	GTPase activating
' '	0,,0,0010,000	.,	1	5,55	0,0 .		protein-like
417	CCCAGAGACCC	2,31	19	8,23	10,13	Hs.21223	calponin 1, basic,
			l	•	•		smooth muscle
418	GCAAGAAAGTG	2,84	23	8,10	12,05	Hs.155376	hemoglobin, beta
419	AAACAATAAAA	0	8	8,00	12,08	Hs.229971	EST
420	CGTGGGACACT	0	. 8	8,00	12,08	Hs.110196	NICE-1 protein
421	CTGTTCTCTTG	0,5	4	8,00	2,33	Hs.46824	ESTs
422	TAGTTGGAAAA	4,16	33	7,93	16,82	Hs.1119	nuclear receptor
							subfamily 4, group A,
							member 1
-	GTGAAAGCCTG	0,76	6	7,89		Hs.258926	EST
424	TTGGTTTGCTG	0,38	3	7,89	1,76	Hs.284326	Human clone 23960
405	00707447704	0.00		7.00	4 70		mRNA sequence
425	CCTGTAATTCA	0,38	3	7,89	1,76	Hs.277331	EST, Weakly similar to
1			- 1				ALU5_HUMAN ALU SUBFAMILY
426	AAACCCCGTCT	0,38	3	7 90	1.76	Hs.273464	ESTs, Weakly similar to
420	AAACCCCGICI	0,36	अ	7,89	1,70	ПS.213464 	ALU2 HUMAN ALU
•							SUBFAMILY
427	ATCGCACTACT	0,38	3	7,89	1 76	Hs.161721	ESTs
	TTGAGGGGGTG	2,17	17	7,83		Hs.76549	(Manual assignment)
		_,		.,00	0,00	10.7 00 10	MEMOREC
	4						unassignable (probably r
429	TTTGGTTTTCC	14,74	115	7,80	56,2	Hs.179573	collagen, type I, alpha 2
430	ACAAAACCCCG	0,91	7	7,69		Hs.259505	EST
431	GTGCTCAATAG	0,26	2	7,69	1,18	Hs.8687	ESTs
432	CAACCAGTAAA	0,26	2	7,69	1,18	Hs.79914	lumican
433	AGTTTATGCCC	0,26	2	7,69		Hs.76591	KIAA0887 protein
	TTCACATTGTC	0,26	2	7,69		Hs.285804	ESTs
	CCTGGCCTAGA	0,26	2	7,69		Hs.285472	ESTs
	AAACTGGGAGG	0,26	2	7,69		Hs.231722	ESTs
	GCCGCCTTCT	0,26	2	7,69		Hs.201292	ESTs
438	TGCAGGTTTGT	0,26	2	7,69	1,18	Hs.183800	Ran GTPase activating
							protein 1
	CTCAACTTGTA	0,26	2	7,69		Hs.117582	CGI-43 protein
	TTTGCTTTTGT	1,32	10	7,58		Hs.234642	aquaporin 3
441	TAAATGAAAAA	0,41	3	7,32	1,69	Hs.82120	nuclear receptor
			1				subfamily 4, group A,
440	CCCCCTTCCT	0.44		7.00	4.00	He 240000	member 2
442	GCCCCCTTCCT	0,41	3	7,32	1,69	Hs.212680	tumor necrosis factor
			İ				receptor superfamily, member 1
443	CCTGTAATTGC	0,41	3	7,32	1 60	Hs.181464	ESTs
	TCACCCTCCAG	0,41	3	7,32		Hs.15251	hypothetical protein
	AAGCTCTGTGT	0,41	5	7,14		Hs.19813	ESTs Protein
	ATGGTGGGCGC	0,56				Hs.266417	EST
1 775	r 55155566	1 2,20	-7	1,17	2,13	J. 10.2007 17	1-01

ALUS   HUMAN ALU   SUBFAMILY	147	CCTGTAGTTCT	0,56	4	7,14	2 10	Hs.231918	ESTs, Weakly similar to
SUBFÄMILY   STS, Weakly similar to prostate-specific transgluta   SUBFAMILY   STS, Weakly similar to prostate-specific transgluta   STS	447	CCIGIAGITOI	0,50	7	7,17	2, 19	115.251910	
448   CATCTGTAATC								
A49   TCTATAATCCC   0,85   6   7,06   3,14   Hs.96866   ESTs	448	CATCTGTAATC	0.56	4	7 14	2 19	Hs 153290	
			0,00	1	• ,	۷, ۱۰	110.100200	
449   TCTATAATCCC								1.
450 ATGGCACGTGC	449	TCTATAATCCC	0,85	6	7,06	3,14	Hs.96866	
451 CTGGTGCACTG	450	ATGGCACGTGC	1	7				stromal cell protein
ALUB_HUMÁN ALU   SUBFAMILY			0,29	2				
452 ATAAAAAGAAA         0,29         2         6,90         1,11 Hs.83942         cathepsin K (pycnodysostosis)           453 CACTTGTAGTC         0,29         2         6,90         1,11 Hs.7845         Homo sapiens cDNA FLJ20820 fis, clone ADSE00490           454 ATTTGGAGAGG         0,29         2         6,90         1,11 Hs.55777         Fukuyama type congenital muscular dystrophy           455 CAGCAGCTTGT         0,29         2         6,90         1,11 Hs.55405         ESTs, Weakly similar to Rab7 [H.sapiens]           456 GTATTTTCATA         0,29         2         6,90         1,11 Hs.29640         suppression of tumorigenicity 15 (reversion-inducing           457 CTTGTTGCAAT         0,29         2         6,90         1,11 Hs.278222         ESTs, Highly similar to endothelial nitric oxide syn           458 CAGGGTGGGTG         0,29         2         6,90         1,11 Hs.249956         EST           460 TGGCATAATCA         0,29         2         6,90         1,11 Hs.297522         ESTs           461 CCTTCCTCCC         0,29         2         6,90         1,11 Hs.199752         ESTs           462 CCCTGAATGAA         0,29         2         6,90         1,11 Hs.199752         ESTs           463 GATTAGCACCA         0,29         2         6,90         1,11 H				1		•		
453   CACTTGTAGTC   0,29   2   6,90   1,11   Hs.7845   Home sapiens cDNA FLJ20820 fis, clone ADSE00490   454   ATTTGGAGAGG   0,29   2   6,90   1,11   Hs.55777   Fukuyama type congenital muscular dystrophy   455   CAGCAGCTTGT   0,29   2   6,90   1,11   Hs.55405   ESTs, Weakly similar to Rab7   H.sapiens   456   GTATTTCATA   0,29   2   6,90   1,11   Hs.42140   hypothetical protein FLJ10103   457   CTTGTTGCAAT   0,29   2   6,90   1,11   Hs.29640   suppression of tumorigenicity 15 (reversion-inducing   458   CAGGGTGGGTG   0,29   2   6,90   1,11   Hs.278222   ESTs, Highly similar to endothelial nitric oxide syn   459   AGCCACTACGC   0,29   2   6,90   1,11   Hs.249956   EST   460   TGGCATAATCA   0,29   2   6,90   1,11   Hs.199752   ESTs   461   CCTTCCTCC   0,29   2   6,90   1,11   Hs.199545   frizzled (Drosophila) homolog 4   463   GATTAGCACA   0,29   2   6,90   1,11   Hs.19545   frizzled (Drosophila) homolog 4   464   GCCGTGAAAAA   0,29   2   6,90   1,11   Hs.164257   ESTs   465   AGAAAGAAGAA   0,29   2   6,90   1,11   Hs.164257   ESTs   466   GCTTCCTCCT   0,73   5   6,85   2,6   Hs.85289   CD34 antigen   467   TITCTTCCTT   0,73   5   6,85   2,6   Hs.283009   tuftelin 1   468   TGCCTGTAGTC   16,23   111   6,84   49,37   Hs.285275   ESTs, Moderately similar to ALU7   HUMAN ALU   469   CCCTCAATCCC   0,88   6   6,82   3,07   Hs.83077   interleukin 18 (interferongamma-inducing factor)   470   ACAACTTTAT   0,88   6   6,82   3,07   Hs.83077   interleukin 18 (interferongamma-inducing factor)   470   ACAACTTTAT   0,88   6   6,82   3,07   Hs.83078   ESTs   471   GCAAACCTAAA   0,44   3   6,82   1,62   Hs.858516   ESTs   473   TTGAATAGTGA   0,59   4   6,78   2,12   Hs.38516   ESTs   475   TTGAATAGTGA   0,59   4   6,78   2,12   Hs.38516   ESTs   475   TTGAATAGTGA   0,59   4   6,78   2,12   Hs.38516   ESTs   475   TTGAATAGTGA   0,59   4   6,								SUBFAMILY
453   CACTTGTAGTC	452	ATAAAAAGAAA	0,29	. 2	6,90	1,11	Hs.83942	cathepsin K
ATTTGGAGAGG								
ADSE00490   ATTTGGAGAGG   0,29   2   6,90   1,11   Hs.55777   Fukuyama type congenital muscular dystrophy	453	CACTTGTAGTC	0,29	2	6,90	1,11	Hs.7845	
454         ATTTGGAGAGG         0,29         2         6,90         1,11         Hs.55777         Fukuyama type congenital muscular dystrophy           455         CAGCAGCTTGT         0,29         2         6,90         1,11         Hs.55405         ESTs, Weakly similar to Rab7 [H.sapiens]           456         GTATTTTCATA         0,29         2         6,90         1,11         Hs.42140         hypothetical protein FLJ10103           457         CTTGTTGCAAT         0,29         2         6,90         1,11         Hs.29640         suppression of tumorigenicity 15 (reversion-inducing           458         CAGGGTGGGTG         0,29         2         6,90         1,11         Hs.249956         EST           459         AGCCACTACGC         0,29         2         6,90         1,11         Hs.249956         EST           460         TGGCATAATCA         0,29         2         6,90         1,11         Hs.19545         ESTs           461         CCTGAATGAA         0,29         2         6,90         1,11         Hs.19545         Firizzled (Drosophila) homolog 4           463         GATTAGCACCA         0,29         2         6,90         1,11         Hs.18046         ribosomal protein L5           464	1			ļ				
Congenital muscular dystrophy   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular to Rab Till Hs.sphile   Congenital muscular dystrophy   Congenital muscu								
455   CAGCAGCTTGT   0,29   2   6,90   1,11   Hs.55405   ESTs, Weakly similar to Rab7 [H.sapiens]     456   GTATTTTCATA   0,29   2   6,90   1,11   Hs.42140   hypothetical protein FLJ10103     457   CTTGTTGCAAT   0,29   2   6,90   1,11   Hs.29640   suppression of tumorigenicity 15 (reversion-inducing ESTs, Highly similar to endothelial nitric oxide syn     458   CAGGGTGGGTG   0,29   2   6,90   1,11   Hs.278222   ESTs, Highly similar to endothelial nitric oxide syn     459   AGCCACTACGC   0,29   2   6,90   1,11   Hs.249956   EST     460   TGGCATAATCA   0,29   2   6,90   1,11   Hs.199752   ESTs     461   CCTTCCTCTCC   0,29   2   6,90   1,11   Hs.199752   ESTs     462   CCCTGAATGAA   0,29   2   6,90   1,11   Hs.199752   ESTs     463   GATTAGCACCA   0,29   2   6,90   1,11   Hs.180946   ribosomal protein L5     464   GCCGTGAAAAA   0,29   2   6,90   1,11   Hs.180946   ribosomal protein L5     465   AGAAAGAAGGA   0,29   2   6,90   1,11   Hs.1501   syndecan 2 (heparan sulfate proteoglycan 1, cell sur     466   GCTTCCTCCTC   0,73   5   6,85   2,6   Hs.283009   tuftelin 1     468   TGCCTGTAGTC   16,23   111   6,84   49,37   Hs.285275   ESTs, Moderately similar to ALUT_HUMAN ALU     469   CCCTCAATCCC   0,88   6   6,82   3,07   Hs.283213   EST     470   ACAACTTTAT   0,88   6   6,82   3,07   Hs.283213   EST     471   GCAAACCTAAA   0,44   3   6,82   1,62   Hs.86686   prefoldin 5     472   GAGTGCAACCC   0,44   3   6,82   1,62   Hs.86686   prefoldin 5     473   TTGAATAGTGA   0,59   4   6,78   2,12   Hs.38516   ESTs	454	ATTTGGAGAGG	0,29	2	6,90	1,11	Hs.55777	
455         CAGCAGCTTGT         0,29         2         6,90         1,11         Hs.55405         ESTs, Weakly similar to Rab7 [H.sapiens]           456         GTATTTCATA         0,29         2         6,90         1,11         Hs.42140         hypothetical protein FLJ10103           457         CTTGTTGCAAT         0,29         2         6,90         1,11         Hs.29640         suppression of tumorigenicity 15 (reversion-inducing           458         CAGGGTGGGTG         0,29         2         6,90         1,11         Hs.278222         ESTs, Highly similar to endothelial nitric oxide syn           459         AGCCACTACGC         0,29         2         6,90         1,11         Hs.249956         EST           460         TGGCATAATCA         0,29         2         6,90         1,11         Hs.237063         ESTs           461         CCTTCCTCCC         0,29         2         6,90         1,11         Hs.199752         ESTs           462         CCCTGAATGAA         0,29         2         6,90         1,11         Hs.199545         frizzled (Drosophila) homolog 4           463         GATTAGCACCA         0,29         2         6,90         1,11         Hs.180457         ESTs           464         <		•		1				
Rab7 [H.sapiens]   Rab7 [H.sapiens]	455	CACCACCTTCT	0.00		0.00	4 4 4	11- 55 405	
456         GTATTTCATA         0,29         2         6,90         1,11         Hs.42140         hypothetical protein FLJ10103           457         CTTGTTGCAAT         0,29         2         6,90         1,11         Hs.29640         suppression of tumorigenicity 15 (reversion-inducing           458         CAGGGTGGGTG         0,29         2         6,90         1,11         Hs.278222         ESTs, Highly similar to endothelial nitric oxide syn           459         AGCCACTACGC         0,29         2         6,90         1,11         Hs.249956         EST           460         TGGCATAATCA         0,29         2         6,90         1,11         Hs.237063         ESTs           461         CCTTCCTCCC         0,29         2         6,90         1,11         Hs.199752         ESTs           462         CCCTGAATGAA         0,29         2         6,90         1,11         Hs.19545         frizzled (Drosophila) homolog 4           463         GATTAGCACCA         0,29         2         6,90         1,11         Hs.180946         ribosomal protein L5           464         GCCGTGAAAAA         0,29         2         6,90         1,11         Hs.190946         ribosomal protein L5           465         A	455	CAGCAGCIIGI	0,29	2	6,90	1,11	HS.554U5	
Section   Comparison   Compar	156	GTATITICATA .	0.20	2	6 90	1 11	He 42140	
457         CTTGTTGCAAT         0,29         2         6,90         1,11         Hs.29640         suppression of tumorigenicity 15 (reversion-inducing           458         CAGGGTGGGTG         0,29         2         6,90         1,11         Hs.278222         ESTs, Highly similar to endothelial nitric oxide syn           459         AGCCACTACGC         0,29         2         6,90         1,11         Hs.237063         EST           460         TGGCATAATCA         0,29         2         6,90         1,11         Hs.237063         ESTs           461         CCTTCCTCCC         0,29         2         6,90         1,11         Hs.199752         ESTs           462         CCCTGAATGAA         0,29         2         6,90         1,11         Hs.199752         ESTs           464         GCCGTGAAAAA         0,29         2         6,90         1,11         Hs.180946         ribosomal protein L5           465         AGAAAGAAGGA         0,29         2         6,90         1,11         Hs.164257         ESTs           466         GCTTCCTCCTC         0,73         5         6,85         2,6         Hs.285289         CD34 antigen           467         TTTCTTCCCTT         0,73         5	430	GIAITITOAIA	0,29	4	0,90	1,11	115.42 140	
tumorigenicity 15   treversion-inducing   tumorigenicity 15   treversion-inducing   test	457	CTTGTTGCAAT	0.29	2	6.90	1 11	Hs 29640	
CAGGGTGGGTG		011011007111	0,20	7	0,00	.,		
458         CAGGGTGGGTG         0,29         2         6,90         1,11         Hs.278222         ESTs, Highly similar to endothelial nitric oxide syn           459         AGCCACTACGC         0,29         2         6,90         1,11         Hs.249956         EST           460         TGGCATAATCA         0,29         2         6,90         1,11         Hs.237063         ESTs           461         CCTTCCTCC         0,29         2         6,90         1,11         Hs.199752         ESTs           462         CCCTGAATGAA         0,29         2         6,90         1,11         Hs.19545         frizzled (Drosophila) homolog 4           463         GATTAGCACCA         0,29         2         6,90         1,11         Hs.180946         ribosomal protein L5           464         GCCGTGAAAAA         0,29         2         6,90         1,11         Hs.164257         ESTs           465         AGAAAGAAGGA         0,29         2         6,90         1,11         Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466         GCTTCCTCCTC         0,73         5         6,85         2,6 Hs.283009         tuftelin 1           469         TCTCTAGTC         16,23         111<				[				
Head of the lian intric oxide   Head of the lian intriceus   Head of the lian intriceus   Head of the lian intriceus   Head of the lian intriceus   Head of the lian intriceus   Head of the lian intriceus   Head of the li	458	CAGGGTGGGTG	0.29	2	6,90	1.11	Hs.278222	
459 AGCCACTACGC         0,29         2         6,90         1,11 Hs.249956         EST           460 TGGCATAATCA         0,29         2         6,90         1,11 Hs.237063         ESTs           461 CCTTCCTCTC         0,29         2         6,90         1,11 Hs.199752         ESTs           462 CCCTGAATGAA         0,29         2         6,90         1,11 Hs.19545         frizzled (Drosophila) homolog 4           463 GATTAGCACCA         0,29         2         6,90         1,11 Hs.180946         ribosomal protein L5           464 GCCGTGAAAAA         0,29         2         6,90         1,11 Hs.164257         ESTs           465 AGAAAGAAGGA         0,29         2         6,90         1,11 Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466 GCTTCCTCCTC         0,73         5         6,85         2,6 Hs.85289         CD34 antigen           467 TTTCTTCCCTT         0,73         5         6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGAGTC         16,23         111         6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6         6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA					·	•		
460 TGGCATAATCA         0,29         2         6,90         1,11 Hs.237063         ESTs           461 CCTTCCTCTC         0,29         2         6,90         1,11 Hs.199752         ESTs           462 CCCTGAATGAA         0,29         2         6,90         1,11 Hs.19545         frizzled (Drosophila) homolog 4           463 GATTAGCACCA         0,29         2         6,90         1,11 Hs.180946         ribosomal protein L5           464 GCCGTGAAAAA         0,29         2         6,90         1,11 Hs.164257         ESTs           465 AGAAAGAAGGA         0,29         2         6,90         1,11 Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466 GCTTCCTCCTC         0,73         5         6,85         2,6 Hs.85289         CD34 antigen           467 TTTCTTCCCTT         0,73         5         6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGTAGTC         16,23         111         6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6         6,82         3,07 Hs.283213         EST           470 ACAACTTTAT         0,88         6         6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA				- 1				syn
461 CCTTCCTCC         0,29         2         6,90         1,11 Hs.199752         ESTs           462 CCCTGAATGAA         0,29         2         6,90         1,11 Hs.19545         frizzled (Drosophila) homolog 4           463 GATTAGCACCA         0,29         2         6,90         1,11 Hs.180946         ribosomal protein L5           464 GCCGTGAAAAA         0,29         2         6,90         1,11 Hs.164257         ESTs           465 AGAAAGAAGGA         0,29         2         6,90         1,11 Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466 GCTTCCTCCTC         0,73         5         6,85         2,6 Hs.85289         CD34 antigen           467 TTTCTCCCTT         0,73         5         6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGTAGTC         16,23         111         6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6         6,82         3,07 Hs.83077         interleukin 18 (interferongamma-inducing factor)           470 ACAACTTTAT         0,88         6         6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA         0,44         3         6,82         1,62 Hs.54680         ESTs <tr< td=""><td>459</td><td>AGCCACTACGC</td><td></td><td>2</td><td>6,90</td><td></td><td></td><td>EST</td></tr<>	459	AGCCACTACGC		2	6,90			EST
462         CCCTGAATGAA         0,29         2         6,90         1,11         Hs.19545         frizzled (Drosophila) homolog 4           463         GATTAGCACCA         0,29         2         6,90         1,11         Hs.180946         ribosomal protein L5           464         GCCGTGAAAAA         0,29         2         6,90         1,11         Hs.164257         ESTs           465         AGAAAGAAGGA         0,29         2         6,90         1,11         Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466         GCTTCCTCCTC         0,73         5         6,85         2,6         Hs.85289         CD34 antigen           467         TTTCTTCCCTT         0,73         5         6,85         2,6         Hs.283009         tuftelin 1           468         TGCCTGTAGTC         16,23         111         6,84         49,37         Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469         CCCTCAATCCC         0,88         6         6,82         3,07         Hs.83077         interleukin 18 (interferongamma-inducing factor)           470         ACAACTTTAT         0,88         6         6,82         3,07         Hs.283213         EST           471			0,29		6,90	1,11	Hs.237063	
homolog 4   463 GATTAGCACCA   0,29   2   6,90   1,11 Hs.180946   ribosomal protein L5   464 GCCGTGAAAAA   0,29   2   6,90   1,11 Hs.164257   ESTs   465 AGAAAGAAGGA   0,29   2   6,90   1,11 Hs.1501   syndecan 2 (heparan sulfate proteoglycan 1, cell sur   466 GCTTCCTCCTC   0,73   5   6,85   2,6 Hs.85289   CD34 antigen   467 TTTCTTCCCTT   0,73   5   6,85   2,6 Hs.283009   tuftelin 1   468 TGCCTGTAGTC   16,23   111   6,84   49,37 Hs.285275   ESTs, Moderately similar to ALU7 HUMAN ALU   469 CCCTCAATCCC   0,88   6   6,82   3,07 Hs.83077   interleukin 18 (interferongamma-inducing factor)   470 ACAACTTTTAT   0,88   6   6,82   3,07 Hs.283213   EST   471 GCAAACCTAAA   0,44   3   6,82   1,62 Hs.80686   prefoldin 5   472 GAGTGCAACCC   0,44   3   6,82   1,62 Hs.54680   ESTs   473 TTGAATAGTGA   0,59   4   6,78   2,12 Hs.38516   ESTs								
463 GATTAGCACCA         0,29         2         6,90         1,11 Hs.180946 ribosomal protein L5           464 GCCGTGAAAAA         0,29         2         6,90         1,11 Hs.164257 ESTs           465 AGAAAGAAGGA         0,29         2         6,90         1,11 Hs.1501 syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466 GCTTCCTCCTC         0,73         5         6,85         2,6 Hs.85289 CD34 antigen           467 TTTCTTCCCTT         0,73         5         6,85         2,6 Hs.283009 fuffelin 1           468 TGCCTGTAGTC         16,23         111 6,84         49,37 Hs.285275 ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6         6,82         3,07 Hs.83077 interleukin 18 (interferongamma-inducing factor)           470 ACAACTTITAT         0,88         6         6,82         3,07 Hs.283213 EST           471 GCAAACCTAAA         0,44         3         6,82         1,62 Hs.80686 prefoldin 5           472 GAGTGCAACCC         0,44         3         6,82         1,62 Hs.54680 ESTs           473 TTGAATAGTGA         0,59         4         6,78         2,12 Hs.38516 ESTs	462	CCCTGAATGAA	0,29	2	6,90	1,11	Hs.19545	
464 GCCGTGAAAAA         0,29         2 6,90         1,11 Hs.164257         ESTs           465 AGAAAGAAGGA         0,29         2 6,90         1,11 Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466 GCTTCCTCCTC         0,73         5 6,85         2,6 Hs.85289         CD34 antigen           467 TTTCTTCCCTT         0,73         5 6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGTAGTC         16,23         111 6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6 6,82         3,07 Hs.83077         interleukin 18 (interferongamma-inducing factor)           470 ACAACTTTTAT         0,88         6 6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA         0,44         3 6,82         1,62 Hs.80686         prefoldin 5           472 GAGTGCAACCC         0,44         3 6,82         1,62 Hs.54680         ESTs           473 TTGAATAGTGA         0,59         4 6,78         2,12 Hs.38516         ESTs								
465 AGAAAGAAGGA         0,29         2 6,90         1,11 Hs.1501         syndecan 2 (heparan sulfate proteoglycan 1, cell sur           466 GCTTCCTCCTC         0,73         5 6,85         2,6 Hs.85289         CD34 antigen           467 TTTCTTCCCTT         0,73         5 6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGTAGTC         16,23         111         6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU interleukin 18 (interferongamma-inducing factor)           469 CCCTCAATCCC         0,88         6 6,82         3,07 Hs.83077         interleukin 18 (interferongamma-inducing factor)           470 ACAACTTTTAT         0,88         6 6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA         0,44         3 6,82         1,62 Hs.80686         prefoldin 5           472 GAGTGCAACCC         0,44         3 6,82         1,62 Hs.54680         ESTs           473 TTGAATAGTGA         0,59         4 6,78         2,12 Hs.38516         ESTs				2				
Sulfate proteoglycan 1, cell sur								
Cell sur	465	AGAAAGAAGGA	0,29	2	6,90	1,11	Hs.1501	syndecan 2 (heparan
466 GCTTCCTCCTC         0,73         5         6,85         2,6 Hs.85289         CD34 antigen           467 TTTCTTCCCTT         0,73         5         6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGTAGTC         16,23         111         6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6         6,82         3,07 Hs.83077         interleukin 18 (interferongamma-inducing factor)           470 ACAACTTTTAT         0,88         6         6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA         0,44         3         6,82         1,62 Hs.80686         prefoldin 5           472 GAGTGCAACCC         0,44         3         6,82         1,62 Hs.54680         ESTs           473 TTGAATAGTGA         0,59         4         6,78         2,12 Hs.38516         ESTs				İ	· ·			sulfate proteoglycan 1,
467 TTTCTTCCCTT         0,73         5         6,85         2,6 Hs.283009         tuftelin 1           468 TGCCTGTAGTC         16,23         111         6,84         49,37 Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU           469 CCCTCAATCCC         0,88         6         6,82         3,07 Hs.83077         interleukin 18 (interferongamma-inducing factor)           470 ACAACTTTTAT         0,88         6         6,82         3,07 Hs.283213         EST           471 GCAAACCTAAA         0,44         3         6,82         1,62 Hs.80686         prefoldin 5           472 GAGTGCAACCC         0,44         3         6,82         1,62 Hs.54680         ESTs           473 TTGAATAGTGA         0,59         4         6,78         2,12 Hs.38516         ESTs	400	COTTOCTOCTO	0.70		0.05		11- 05000	
468 TGCCTGTAGTC         16,23         111         6,84         49,37         Hs.285275         ESTs, Moderately similar to ALU7_HUMAN ALU to ALU7_HUMAN ALU interleukin 18 (interferongamma-inducing factor)           469 CCCTCAATCCC         0,88         6         6,82         3,07         Hs.83077         interleukin 18 (interferongamma-inducing factor)           470 ACAACTTTAT         0,88         6         6,82         3,07         Hs.283213         EST           471 GCAAACCTAAA         0,44         3         6,82         1,62         Hs.80686         prefoldin 5           472 GAGTGCAACCC         0,44         3         6,82         1,62         Hs.54680         ESTs           473 TTGAATAGTGA         0,59         4         6,78         2,12         Hs.38516         ESTs								
to ALU7_HUMAN ALU  469 CCCTCAATCCC 0,88 6 6,82 3,07 Hs.83077 interleukin 18 (interferongamma-inducing factor)  470 ACAACTTTAT 0,88 6 6,82 3,07 Hs.283213 EST  471 GCAAACCTAAA 0,44 3 6,82 1,62 Hs.80686 prefoldin 5  472 GAGTGCAACCC 0,44 3 6,82 1,62 Hs.54680 ESTs  473 TTGAATAGTGA 0,59 4 6,78 2,12 Hs.38516 ESTs								
469 CCCTCAATCCC       0,88       6       6,82       3,07       Hs.83077       interleukin 18 (interferongamma-inducing factor)         470 ACAACTTTTAT       0,88       6       6,82       3,07       Hs.283213       EST         471 GCAAACCTAAA       0,44       3       6,82       1,62       Hs.80686       prefoldin 5         472 GAGTGCAACCC       0,44       3       6,82       1,62       Hs.54680       ESTs         473 TTGAATAGTGA       0,59       4       6,78       2,12       Hs.38516       ESTs	400	IGOCIGIAGIO	10,23	111	0,64	49,37	IU2.50272	
	460	CCCTCAATCCC	0.88	- 6	6 82	3.07	He 83077	
470 ACAACTTTTAT       0,88       6       6,82       3,07 Hs.283213       EST         471 GCAAACCTAAA       0,44       3       6,82       1,62 Hs.80686       prefoldin 5         472 GAGTGCAACCC       0,44       3       6,82       1,62 Hs.54680       ESTs         473 TTGAATAGTGA       0,59       4       6,78       2,12 Hs.38516       ESTs	703	00010/1000	0,00	쒸	0,02	3,07	13.03077	
471 GCAAACCTAAA       0,44       3 6,82 1,62 Hs.80686 prefoldin 5         472 GAGTGCAACCC       0,44       3 6,82 1,62 Hs.54680 ESTs         473 TTGAATAGTGA       0,59 4 6,78 2,12 Hs.38516 ESTs	470	ACAACTTTAT	0.88	6	6.82	3.07	Hs 283213	
472 GAGTGCAACCC       0,44       3       6,82       1,62 Hs.54680       ESTs         473 TTGAATAGTGA       0,59       4       6,78       2,12 Hs.38516       ESTs								
473 TTGAATAGTGA 0,59 4 6,78 2,12 Hs.38516 ESTs								

475	GCAAAACACTG	0,59	4	6,78	2,12	Hs.198552	Homo sapiens mRNA; cDNA DKFZp566B193 (from clone
476	CATTTGGGAAG	0,59	4	6,78	2,12	Hs.111334	ferritin, light polypeptide
	CAAGAGATGCT	0,15	1	6,67		Hs.99741	ESTs, Weakly similar to cell division control relate
478	AGTTTATTTCA	0,15	1	6,67		Hs.99016	Human DNA sequence from clone 310J6 on chromosome
479	GAGGATCTGCG	0,15	1	6,67	0,54	Hs.90998	KIAA0128 protein; septin
480	ACTCTGGCTCA	0,15	1	6,67	·	Hs.88974	cytochrome b-245, beta polypeptide (chronic granulom
481	GGGGTTAGGGG	0,15	_ 1	6,67	0,54	Hs.85050	phospholamban
482	TAATATATCTG	0,15	1	6,67		Hs.8203	endomembrane protein emp70 precursor isolog
483	TAAGTCTATAT	0,15	1	6,67	0,54	Hs.78864	Fc fragment of IgG, low affinity IIa, receptor for (
484	CACAAAAGGAT	0,15	1	6,67	0,54	Hs.77603	ESTs
485	GAGAAACCCTT	0,15	1	6,67	0,54	Hs.7739	ESTs
486	GAAAACAGTAA	0,15	1	6,67	0,54	Hs.76111	dystroglycan 1 (dystrophin-associated glycoprotein 1
487	GGCAATTTACT	0,15	1	6,67	0,54	Hs.75813	polycystic kidney disease 1 (autosomal dominant)
	TTTTGTGCTAG	0,15	1	6,67		Hs.74649	cytochrome c oxidase subunit VIc
	TTCCCGTGGCT	0,15	1	6,67	· .	Hs.70983	PTPL1-associated . RhoGAP 1
	GGCCCAGGCCT	0,15	. 1	6,67		Hs.575	aldehyde dehydrogenase 3
-	CGGCCACGTAT	0,15	. 1	6,67		Hs.55993	ESTs
492	TATAGCTGCAT	0,15	1	6,67	0,54	Hs.55964	ESTs, Weakly similar to C4HU complement C4A precurso
493	CTTGTGTTTAT	0,15	1	6,67	0,54	Hs:50748	chromosome 21 open reading frame 18
$\overline{}$	CACAAAAATGC	0,15	1	6,67		Hs.49944	ESTs
	CTGTAATTTTA	0,15	1	6,67	0,54	Hs.48480	ESTs
	CAATTCTTTCT	0,15	1	6,67	0,54	Hs.48403	hypothetical protein FLJ10847
497	ATTCTGCAGAG	0,15	1	6,67	0,54	Hs.47232	ESTs
	TTTATATCATT	0,15	1	6,67		Hs.47099	ESTs
	ACTCCATAAAA	0,15	1	6,67	0,54	Hs.4273	Human DNA sequence from clone RP1-104A17 on
500	TTTTCATTATA	0,15	1	6,67	0,54	Hs.42656	ESTs, Moderately similar to ALU1_HUMAN ALU

		=		=1			1
	GACTCGACCAG	0,15	1	6,67		Hs.31922	ESTs
	GGTGTGTTTTA	0,15	1	6,67		Hs.31566	ESTs
	ATAAATTTATG	0,15	1	6,67		Hs.30715	ESTs
504	GGAGGCCGAGA	0,15	1	6,67		Hs.285565	ESTs
505	ATATTCAGCTG	0,15	1	6,67	0,54	Hs.285379	ESTs
506	GAACTTGTCTG	0,15	1	6,67	0,54	Hs.279934	Homo sapiens mRNA;
				1			cDNA DKFZp434I0835
							(from clone
507	CCCGTATATGT	0,15	1	6,67	0,54	Hs.279844	hypothetical protein
		· .				· · · · · · · · · · · · · · · · · · ·	FLJ10033
	GTGGACCTGAG	0,15	1	6,67		Hs.279059	ESTs
-	AGCCTGGAAGG_	0,15	1	6,67		Hs.278549	ESTs
	TGGTTTTTGAG	0,15	1	6,67		Hs.275865	ribosomal protein S18
·511	GCTTGTTCAAA	0,15	1	6,67	0,54	Hs.274969	ESTs
512	TCCACCAGCCA	0,15	1	6,67	0,54	Hs.27457	ESTs
513	GCACTCCAACC	0,15	1	6,67		Hs.273682	EST
514	CCCATAATCCT	0,15	1	6,67	0,54	Hs.270797	ESTs
515	AGTCACAGCTT	0,15	1	6,67	0,54	Hs.267448	hypothetical protein
							FLJ20039
516	TAGAAAAAACC	0,15	1	6,67	0,54	Hs.262476	S-adenosylmethionine
							decarboxylase 1
517	TAAAGATGGCA	0,15	1	6,67	0,54	Hs.25357	Homo sapiens clone
• × •							24488 mRNA sequence
518	GGCTATGCCCT	0,15	1	6,67		Hs.243855	ESTs
519	AGACATTGACA	0,15	. 1	6,67		Hs.239934	CGI-96 protein
520	TATGCTTTAAA	0,15	1	6,67	0,54	Hs.237225	ribosomal protein S5
							pseudogene 1
521	TATTGCTAAAT	0,15	1	6,67	0,54	Hs.23590	solute carrier family 16
	<sub>.</sub>				٠. '	1 1	(monocarboxylic acid
		·				~	transp
	CCAAGGCACTG	0,15	1	6,67		Hs.234863	ESTs
	AACTTTCCAAA	0,15	1	6,67		Hs.23457	ESTs
	TCTCACAAGGG	0,15	1	6,67		Hs.233476	ESTs
	CCACTATGCCT	0,15	1	6,67		Hs.231229	ESTs
	GGGCGCCTGGC	0,15	1	6,67		Hs.224242	EST
527	GGGGGAAAAA	0,15	1	6,67	0,54	Hs.223590	EST, weakly similar to
			ļ		-		RL3_HUMAN 60S
ļ							RIBOSOMAL
528	TTGTTTATGTA	0,15	1	6,67	0,54	Hs.21958	Homo sapiens cDNA
			ļ				FLJ10532 fis, clone
L							NT2RP2001044
	TTGAGATAAGA	0,15	1	6,67		Hs.21887	ESTs
	GCTATATCCAA	0,15	1	6,67		Hs.218008	ESTs
	TTTGGTTTTCT	0,15	1	6,67		Hs.21431	suppressor of fused
	GATTGTCCTTG	0,15	1	6,67		Hs.211517	ESTs
	GGTTATCAAAG	0,15	1	6,67		Hs.208334	ESTs
	GTCTCGCTGAC	0,15	1	6,67		Hs.207911	ESTs
	AGCCATCGCGC	0,15	1	6,67		Hs.207749	EST
1 526	ACCACCCGTGT	0,15	1	6,67	0.54	Hs.202033	EST

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<u> </u>	A CA OTOTTOOT	0.45		2 2 2	0.54		TEOT
	ACACTCTTCCT	0,15	1	6,67		Hs.20103	ESTs
	GGAGATGTTTG	0,15	1	6,67		Hs.199545	ESTs
	CTCAACCTTAA	0,15	1	6,67		Hs.199243	KIAA0231 protein
	GCAAGACTCCC	0,15	1	6,67		Hs.198011	ESTs
541	CAATACTATTC	0,15	1	6,67	0,54	Hs.197642	hypothetical protein FLJ10388
542	GTTGTGCTCAG	0,15	1	6,67	0,54	Hs.191228	hypothetical protein FLJ20284
543	GGGCTAGCACT	0,15	1	6,67	0,54	Hs.190722	ESTs
544	ACATTCACGCC	0,15	1	6,67	0,54	Hs.184141	glutaryl-Coenzyme A dehydrogenase
545	TATATTTAGTT	0,15	1	6,67	0,54	Hs.183037	protein kinase, cAMP- dependent, regulatory, type I,
546	CCAGCTAGCGA	0,15	1	6,67	0,54	Hs.179756	LW-1
547	TGAGGATACAG	0,15	1	6,67		Hs.177820	Homo sapiens mRNA; cDNA DKFZp564L102 (from clone
	TGCCTACAGTC	0,15	1	6,67		Hs.176207	EST
549	CTCATATGCAA	0,15	1	6,67	0,54	Hs.173540	ATPase, Class V, type 10D
550	AAGGAGGTGGA	0,15	1	6,67	0,54	Hs.172730	ESTs
551	CAAGCCCTGCC	0,15	1	6,67		Hs.172035	hypothetical protein similar to mouse HN1 (Hematolo
552	GCGCTGGGAGG	0,15	1	6,67	0,54	Hs.171763	CD22 antigen
553	TCCTTGGCGTG	0,15	1	6,67	0,54	Hs.168640	Ank, mouse, homolog of
554	TTGCCATATGC	0,15	1	6,67	0,54	Hs.164024	ESTs, Weakly similar to unnamed protein product [H.s
555	GCTCCCTTCAC	0,15	1	6,67	0,54	Hs.162222	EST
556	CTGTGCAAGGA	0,15	1	6,67	0,54	Hs.161554	hypothetical protein FLJ20159
	TGTGCTTGTGT	0,15	1	6,67		Hs.161554	hypothetical protein FLJ20159
	СТТТТССССС	0,15	1	6,67	· · · · ·	Hs.156007	Down syndrome critical region gene 1-like 1
	TCTTTTCTTT	0,15	. 1	6,67		Hs.155606	paired mesoderm homeo box 1
	AAAAGCTGTTT	0,15	1	6,67		Hs.15550	ESTs
	GTCCCCCCCC	0,15	1	6,67		Hs.152454	ESTs
	CCTGGCGGGAT	0,15	1	6,67		Hs.149347	ESTs, Highly similar to KIAA1043 protein [H.sapiens]
	GATTGGTATGA	0,15	1	6,67		Hs.147049	cut (Drosophila)-like 1 (CCAAT displacement protein)
	GTGAAGACTAC	0,15	1	6,67		Hs.14665	ESTs
565	TAAATGGGTTG	0,15	1	6,67	0,54	Hs.1395	early growth response 2 (Krox-20 (Drosophila)

	·				, ——		<del></del>
							homolo
	TGTAAACTTTG	0,15		6,67		Hs.13849	ESTs
567	GACACGTTGCC	0,15	1	6,67	0,54	Hs.136574	arachidonate 12-
							lipoxygenase, 12R type
	TGTGAATTTTA	0,15		6,67		Hs.132834	hematopoietic protein 1
	TAAAACACTTG	0,15		6,67		Hs.130636	ESTs
	TGGCAATTTTC	0,15		6,67		Hs.129636	ESTs
571	TCTGTAGCACA	0,15	1	6,67	0,54	Hs.128766	hypothetical protein FLJ10600
572	TCAGCAGTTAA	0,15	1	6,67	0,54	Hs.128571	ESTs
573	AATGAGGTGCT	0,15	1	6,67	0,54	Hs.128400	ESTs
574	TTCAGAATCTT	0,15	1	6,67	0,54	Hs.125914	ESTs
575	GAATAAGATAT	0,15	1	6,67	0,54	Hs.12479	associated molecule with the SH3 domain of STAM
576	TGTCCCAGCCA	0,15	1	6,67	0,54	Hs.1211	acid phosphatase 5, tartrate resistant
577	CCCATCTCAGA	0,15	1	6,67	0,54	Hs.120769	Homo sapiens cDNA FLJ20463 fis, clone KAT06143
578	AGAAGCTCCAG	0,15	. 1	6,67	0,54	Hs.120021	DKFZP434l092 protein
579	AGTCTTCAAAA	0,15	1	6,67	0,54	Hs.117582	CGI-43 protein
580	GAGTTTTCATT	0,15	1	6,67		Hs.117582	CGI-43 protein
581	TTTTGCAATAA	0,15	1	6,67		Hs.11614	ESTs
582	AGCTCTATGAG	0,15	1	6,67		Hs.115831	ESTs
583	AATAGGTCCCA	0,15	1	6,67		Hs.113029	ribosomal protein S25
584	TAGTCTACTGT	0,15	1	6,67		Hs.112472	ESTs
585	ATCCTTACATC	0,15	1	6,67		Hs.111720	ESTs
586	AACTTTCATAT	0,15	1	6,67		Hs.108787	phosphatidylinositol glycan, class N
587	CCGTAAAAAAA	0,15	. 1	6,67	0,54	Hs.107187	divalent cation tolerant protein CUTA
	TGAAGATGTAA	0,15	1	6,67	0,54	Hs.106217	ESTs
	GAACCATTTGC	0,15		6,67		Hs.104305	KIAA0926 protein
	CTGTGGAGCTG	0,15		6,67	0,54	Hs.103379	ESTs
	AGAATTACAGA	0,15	1	6,67		Hs.101915	ESTs
	TGTTGCTCCCA	0,76	5		2,54	Hs.82210	zinc finger protein 220
	TAGTTTGAAGG	0,76	5	6,58	2,54	Hs.79033	glutaminyl-peptide cyclotransferase (glutaminyl cycl
	CCCAACGCGCT	7,18				Hs.272572	hemoglobin, alpha 2
	CCTATAATCTC	1,99				Hs.117582	CGI-43 protein
	TCTCCTGGACT	0,62	4			Hs.50915	kallikrein 5
597	ATCTTGCCACT	0,62	4	6,45	2,06	Hs.32945	glutamate receptor, metabotropic 1
	AGGATAAAAAA	0,47	3	6,38	1,56	Hs.79404	neuron-specific protein
	GACCACAAATA	0,47		6,38		Hs.76476	cathepsin H
	AGCATATCTTC	0,47	3	6,38	1,56	Hs.275865	ribosomal protein S18
601	AACACAGGAGG	0,47			1,56	Hs.222874	ESTs, Moderately similar

		г			<u> </u>		T
ł			1				to zinc transporter 4
	00000100100	0.47			4.50		[H.sa
602	CGGGGACGAGG	0,47	3	6,38	1,56	Hs.124942	protein phosphatase 2A
		1 1					48 kDa regulatory
<u> </u>							subunit
603	GCGAAGCCCCG	1,11	7	6,31	3,35	Hs.103804	heterogeneous nuclear
			1				ribonucleoprotein U
L	<u></u>	ļ					(scaffold
	GTGGCAGGTAC	1,44	9	6,25		Hs.190467	EST
605	TCTGTTGTTCA	1,44	9	6,25	4,21	Hs.117582	CGI-43 protein
-	CCTGTATCCCA	0,64	4	6,25		Hs.270072	ESTs
607	ATGGATGCTTG	0,32	2	6,25	1,05	Hs.89404	msh (Drosophila) homeo
	·						box homolog 2
608	GACTTCTGTCC	0,32	2	6,25	1,05	Hs.87539	aldehyde
							dehydrogenase 8
609	AAAAAGAAACT	0,32	2	6,25	1,05	Hs.73287	KIAA1235 protein
610	TACTGAAAAAA	0,32	2	6,25	1,05	Hs.5111	hypothetical protein
				**			FLJ20729
611	CACCTGGAGGC	0,32	. 2	6,25	1,05	Hs.30864	ESTs
	GCAAGAGCCCA	0,32	2	6,25		Hs.26670	Human PAC clone RP3-
				,	.,		515N1 from 22q11.2-q22
613	AACCCGGGGAG	0,32	2	6,25	1.05	Hs.228009	EST
	TTGCCCAGGGT	0,32	2	6,25		Hs.225093	ESTs, Weakly similar to
	- 1	0,0-	7	ا - , - ا	.,		ALU5 HUMAN ALU
	· *(-, -, -, -, -, -, -, -, -, -, -, -, -, -	· . ·					SUBFAMILY
615	AAGCAGTTACA	0,32	2	6,25	1.05	Hs.22116	CDC14 (cell division
		-,		-,	.,		cycle 14, S. cerevisiae)
			[	[			homolo
616	GCAGTCATACA	0,32	2	6,25	1,05	Hs.182626	chromosome 22 open
		· 1			•		reading frame 5
617	GGGCACACAC	0,32	2	6,25	1.05	Hs.181900	ESTs
	AGGGAAGGTGA	0,32	2	6,25		Hs.126927	ESTs .
	AGCCGCTGTGC	0,32	2	6,25		Hs.106771	ESTs
	TCAAGCCATCA	5,68	35	6,16		Hs.738	early growth response 1
	GTGGCGGGCAT	0,82	5	6,10		Hs.230564	EST
	ACCTGGGTGCT	0,82	5	6,10		Hs.159643	ESTs, Weakly similar to
		0,02	7	0, 10	-, '-	110.100040	MLD [H.sapiens]
623	TGTGGCGTATA	1,32	8	6,06	3.60	Hs.211582	myosin, light polypeptide
020	10100001/11/1	1,02	٩	0,00	5,05	113.211302	lkinase
624	CGGGCACCTTC	0	6	6,00	8 00	Hs.198249	gap junction protein,
024	000000000000000000000000000000000000000	٦	٩	0,00	0,55	113.130243	beta 5 (connexin 31.1)
625	CCCTTGAGGAG	0	6	6.00	9.00	Hs.1076	
023	COCTIONGONG	١	٥	6,00	0,59	1 15.1076	(Manual assignment)
626	AATGTTTTAA	0.5	3	6.00	1 E	Hs.75335	SPRR1B, cornifin B
1 020		0,5	3	6,00	1,5	118.70000	glycine
							amidinotransferase (L-
		0.5	3	6,00	4 =	Hs.177533	arginine:glycine amidi Homo sapiens mRNA,
E') /	こうしょうしゅう マン・コード			n (III)	1.5	I⊓S. I / / 533	IDOMO SADIENS MKNA
627	CCTCTCCCATT	0,5	ગ	0,00	.,0		
627	CCTCTCCCATT	0,5	1	0,00	.,0		chromosome 1 specific
	TGTTCTGATTT	0,5		6,00		Hs.167835	

	· · · · · ·						oxidase 1, palmitoyl
629	CCTGTAGTGCC	0,85	5	5,88	2.36	Hs.269645	Homo sapiens cell-line
	00101/101000	0,00	٦	0,00	2,00	1.0.2000 10	E8CASS clone E24L
	·		l				estradiol-i
630	CCACTGTATTC	1,03	6	5,83	2.77	Hs.235041	EST
	AGAAATGTATG	1,03	6	5,83		Hs.232068	transcription factor 8
	,	,,,,	. 1	-,	_,		(represses interleukin 2
							expr
632	AGGTCAGAAGA	1,73	10	5,78	4,38	Hs.23437	Homo sapiens mRNA;
			.	1			cDNA DKFZp586G0623
						· · ·	(from clone
633	ATTAAGAAAAT	2,43	14	5,76	5,95	Hs.76549	AHNAK nucleoprotein
							(desmoyokin)
	AACCCGGGGGG	1,05	6	5,71		Hs.6214	KIAA0731 protein
	CACCACAACAA	1,05	6	5,71		Hs.174139	chloride channel 3
$\overline{}$	TTCCCCAGGGT	0,35	2	5,71		Hs.59545	ring finger protein 15
637	CACACTATAGG	0,35	2	5,71	0,99	Hs.58924	ESTs, Weakly similar to
	•						JC5594 jerky gene
							protein ho
	GGGAAAGAGGG	0,35	2	5,71		Hs.35096	KIAA1538 protein
	GCTGGGCGCGG	0,35	2	5,71		Hs.278070	EST
640	GGGGCAACAGC	0,35	2	5,71	0,99	Hs.276770	CDW52 antigen
044	0700010000	0.05		1	0.00	11 000105	(CAMPATH-1 antigen)
-	GTGGCAGGCCC	0,35	2	5,71		Hs.266105	EST
	CATATCCCCTC	0,35	2	5,71		Hs.250746	ESTs
	CTTAGGAGTCA	0,35	2	5,71		Hs.23853	ESTs
644	CAGCACAGTGG	0,35	2	5,71	0,99	Hs.227806	ras GTPase activating
GAE	ATGCTCAAAGG	0,35	2	5,71	0.00	Hs.226018	protein-like Homo sapiens mRNA full
043	AIGCICAAAGG	0,33	4	5,71	0,99	IDS.220010	length insert cDNA clone
	•		.				EURO
646	TTCTGTGCATA	0,35	2	5,71	0.99	Hs.16803	hypothetical protein
0.0	11010100/1//	0,00	-)	١, ا	0,00	1.0.10000	FLJ10231
647	GCATAATGTTT	0,35	2	5,71	0.99	Hs.11050	F-box only protein 9
	TGGCCAGCTCC	3,34	19	5,69		Hs.170121	protein tyrosine
		', '		,,,,,	.,		phosphatase, receptor
				ĺ		•	type, C
649	TCTGGCCCAGC	0,53	3	5,66	1,45	Hs.183	Duffy blood group
650	ATCCTGAGTTA	2,14	12	5,61		Hs.73931	major histocompatibility
		Ì					complex, class II, DQ
							beta
	ACAAATGAAAA	0,18	1	5,56		Hs.96657	hyothetical protein
652	TACATTTCAAG	0,18	1	5,56	0,49	Hs.94376	proprotein convertase
							subtilisin/kexin type 5
653	GCAAAATGCTG	0,18	1	5,56	0,49	Hs.92254	hypothetical protein
25.4							FLJ20163
	TTTAAGAAATG	0,18	1	5,56		Hs.91139	ESTs
655	ACCCACCTGTG	0,18	1	5,56	υ,49	Hs.8736	Homo sapiens mRNA;
						1	cDNA DKFZp564H203
l	I	i l	1			1	(from clone

656	GCAGCAGTGTC	0 10	41	5 56	0.40	Hs.86538	ESTs
		0,18	1	5,56			ESTS
	CCATTAAAAAA	0,18	1	5,56		Hs.85885	
658	AAAGAGGGACG	0,18	1	5,56	0,49	Hs.84229	splicing factor,
				1			arginine/serine-rich 8
050	ACACTTAAAAA	0.40		- F F C	0.40	11- 02204	(suppressor-
009	ACACTTAAAAA	0,18	1	5,56	0,49	Hs.83381	guanine nucleotide
660	TOACAAAAAA	0.40		F 50	0.40	Un 7076	binding protein 11
	TCACAAAAAAA	0,18	1	5,56		Hs.7976	KIAA0332 protein
	ATTTAATATAT	0,18	1	5,56		Hs.7972	KIAA0871 protein
662	AGTTCAAGGCT	0,18	1	5,56	0,49	Hs.79058	suppressor of Ty
							(S.cerevisiae) 4 homolog
000	CACTTOOTATT	0.40		5.50	0.40	11- 70575	
663	GAGTTGCTATT	0,18	1	5,56	0,49	Hs.78575	prosaposin (variant
						•	Gaucher disease and
604	CACTOTOAAA	0.40		F 50	0.40	11- 00000	variant meta
	GACTGTCAAAA	0,18	1	5,56		Hs.69330	ESTs
	GCGCAACAACT	0,18	1	5,56		Hs.61950	DKFZp434A0131 protein
900	TAAATCAGAGA	0,18	7	5,56	0,49	Hs.61929	Homo sapiens cDNA
·							FLJ11010 fis, clone
667	TA A A TO A A TO A	0.40		F 50	0.40	Un 57007	PLACE1003145
$\overline{}$	TAAATGAATGA	0,18	1	5,56		Hs.57967	ESTs
	AATAGGAAGAT	0,18	1	5,56		Hs.50745	ESTs
$\blacksquare$	CAATTAAAAAT	0,18	1	5,56		Hs.50123	zinc finger protein 189
6/0	GTGTATGTGGT	0,18	1	5,56	0,49	Hs.49881	fatty acid binding protein
			ĺ				3, muscle and heart
671	ACAAACCCACA	0,18	1	F FC	0.40	H= 40000	(mamm
67 1	ACAAACCCACA	0,18	1	5,56	0,49	Hs.49282	hypothetical protein
672	CAAGTGAAAGG	0,18	-1	5,56	0.40	Hs.47822	FLJ11088 KIAA0380 gene product;
0/2	CAAGIGAAAGG	0,10		. 5,56	0,49	ПS.47022	RhoA-specific guanine
				l		•	nucleot
673	GAACAGAACGC	0,18	1	5,56	0.49	Hs.47566	ESTs
<b>———</b>	CTAATGGCCCT	0,18	1	5,56		Hs.3416	adipose differentiation-
0,4	01241000001	0,10	'	3,30	0,43	115.5410	related protein
675	TAATGTTTTT	0,18	1	5,56	0.49	Hs.31930	ESTs ESTs
-	ATTTGGCCTGT	0,18	1	5,56		Hs.285519	Homo sapiens OVN6-2
570	ATTIGGCCIGI	0,10	'1	3,30	0,43	1 15.2055 19	mRNA, partial cds
677	TCTGGGAGGGG	0,18	1	5,56	0.49	Hs.285313	core promoter element
"	1010000000	0,10	'1	3,30	0,43	115.200010	binding protein
678	GACTAAATTGT	0,18	1	5,56	0.40	Hs.283643	ESTs .
	TATTITCAÇAA	0,18	1	5,56		Hs.279896	hypothetical protein
0,3	IATTI TOAOAA	0,10	"[	3,30	<i>∪,</i> ~ <i>9</i>	1 13.21 3030	FLJ20546
680	GAGGTTTTCTG	0,18	1	5,56	0.40	Hs.279639	Homo sapiens mRNA;
330		0,10	'	5,50	0,43	13.213033	cDNA DKFZp586M2022
							(from clone
681	CCTAGAATCCC	0,18	1	5,56	0.40	Hs.278982	PRO1779 protein
	CAGCCCCTGTC	0,18	1	5,56		Hs.278234	Homo sapiens mRNA;
552	5.1000001010	3,10	'	5,55	0,73	1.13.27.0204	cDNA DKFZp434H1323
							(from clone
683	GTGTCGGGCTC	0,18	1	5,56	0.40	Hs.278010	EST
1 303	10.01000010	, 5, 10	' '	ارد,	U,73	11 13.27 00 10	1-01

	·						
684	CACCTGTAAAC	0,18	1	5,56	0,49	Hs.277311	EST
685	GCGGCGACTGC	0,18	1	5,56	0,49	Hs.271980	mitogen-activated
			1				protein kinase 6
686	GGAATGAATGA	0,18	1	5,56	0,49	Hs.270824	ESTs
	TGGGATATAGT	0,18	1	5,56	0,49	Hs.269888	ESTs
	GTGGTTGATGC	0,18	1	5,56		Hs.267083	ESTs, Weakly similar to
		-,	ŀ	-,	,		ALU7_HUMAN ALU
			ĺ	1			SUBFAMILY
689	GTGGTAGGTAC	0,18	1	5,56	0.49	Hs.264844	EST, Moderately similar
		0,.0		5,55	٥, ٠٠		to ALU1_HUMAN ALU
690	GCCTGCCTTTA	0,18	1	5,56	0.49	Hs.25371	ESTs, Weakly similar to
		0,.0	1	-,	٥, ٠٠		A37232 mucin, tracheal
.							[H.sa
691	GATCTCGCTTT	0,18	1	5,56	0.49	Hs.250773	signal sequence
00.	0,1101000111	0, 10		0,00	0,0	1.0.2007.10	receptor, alpha
			ļ				(translocon-associat
692	CTCCTATTTTT	0,18	1	5,56	0.49	Hs.25010	hypothetical protein P15-
002		0,10	ï	0,00	0, .0	110.20010	2
693	GTGCGCACCTG	0,18	1	5,56	0.49	Hs.243957	ESTs, Weakly similar to
000	01000000010	0,10	'	5,50	0,40	113.240001	ALU1_HUMAN ALU
			[				SUBFAMILY
694	ATAAAATGTCT	0,18	1	5,56	0.49	Hs.24181	ESTs
	TTTCACCCCGT	0,18	1	5,56		Hs.23492	ESTs
	TATGTTTAAAA	0,18	1	5,56		Hs.234392	platelet-activating factor
030		0,10	'[	3,30	0,43	1 13.207032	acetylhydrolase 2 (40kD)
697	GATCACTTGAG	0,18	1	5,56	0.49	Hs.231798	EST (40KB)
	TTGAGAAAAAA	0,18	1	5,56		Hs.22971	ESTs
	ACCCTGGGAGG	0,18	1	5,56		Hs.228529	ESTs
	TGTGGTGGTGC	0,18	1	5,56		Hs.223618	EST
	GATTGGCCAGA					Hs.21739	
701	GATTOGCCAGA	0,18	'	5,56	0,49	ns.21739	Homo sapiens mRNA;
- 1		ļ	1	1			cDNA DKFZp586I1518
702	COTOACTTATT	0.49		F FC	- 40	11- 046060	(from clone
	GCTGAGTTATT	0,18	1	5,56		Hs.216363	ESTs
703	AGTATTCCTAA	0,18	1	5,56	0,49	Hs.21621	hypothetical protein
704	TOTOTOGOATT	0.40		5.50	0.40	11-04500	DKFZp762O076
	TGTCTGCCATT	0,18		5,56		Hs.21580	ESTs
705	CCCCTGTACTC	0,18	1	5,56	0,49	Hs.213016	ESTs, Highly similar to
							cytokine receptor related
700	07777744404	0.40					pr
	CTTTTTAAAGA	0,18	1	5,56		Hs.212788	EST
707	TACCCGAAAAC	0,18	1	5,56	0,49	Hs.210858	ESTs, Weakly similar to
i			1	1	-		ALUD_HUMAN !!!! ALU
							CLASS D
708	TGCCTCTAGTC	0,18	1	5,56	0,49	Hs.209413	ESTs, Weakly similar to
			.				ALU1_HUMAN ALU
							SUBFAMILY
	CCGCCGCACTC	0,18	1	5,56		Hs.204049	ESTs
710	CTAAGAGGGGT	0,18	1	5,56		Hs.196102	ESTs
711	CTGCTGTAGTC GGACCCTCATT	0,18	1	5,56	0,49	Hs.194319	ESTs

713	AACTAATTCTC	0,18	1	5,56	0.49	Hs.189513	hypothetical protein
							FLJ10213
714	GCAGTGCCAAG	0,18	1	5,56	0,49	Hs.183528	ESTs, Weakly similar to
			}	· }			Bem46-like protein [D.melano
715	ATTCCACCCCA	0.10		5 56	0.40	Hs.180433	rTS beta protein
	ATTGGAGCGCA TGAAAACTCCC	0,18 0,18	1 1	5,56 5,56		Hs.180248	zinc finger protein 124
							(HZF-16)
717	CTCCCAGCCAC	0,18	1	5,56	0,49	Hs.177582	surfactant, pulmonary- associated protein A1
718	GTTTTAAAAAA	0,18	1	5,56	0,49	Hs.174756	EST
719	GAATCAAGCTG	0,18	1	5,56	0,49	Hs.17296	ESTs
720	CTGGGCATTTT	0,18	1	5,56	0,49	Hs.172207	non-POU-domain- containing, octamer- binding
721	TTGTTTGTGTA	0,18	1	5,56		Hs.170162	KIAA1357 protein
	AGGACTGGCAT	0,18	1	5,56		Hs.170114	KIAA0061 protein
	TCGCTTTTAAG	0,18	1	5,56		Hs.168640	Ank, mouse, homolog of
J	AAGTGAAAAAA	0,18	1	5,56		Hs.168159	apoptosis regulator
725	CAAATTAGAAT	0,18	1	5,56	0,49	Hs.162271	ESTs, Moderately similar to CO3_HUMAN COMPLEMENT
726	ATACCACTAAG	0,18	1	5,56	0,49	Hs.153792	5-methyltetrahydrofolate-
-							homocysteine methyltransfer
727	GGTGATGGAGG	0,18	1	5,56	0,49	Hs.149692	ESTs, Highly similar to
				· 1	·		G43284 zinc finger
			1				protein Z
728	TTGGGTTTCTG	0,18	1	5,56	0,49	Hs.147975	ESTs
729	AATAAATGCCC	0,18	1	5,56	0,49	Hs.145522	ESTs
730	CATACACACAT	0,18	1	5,56	0,49	Hs.143460	protein kinase C, nu
731	GTTATACAACA	0,18	1	5,56	0,49	Hs.141183	ESTs
732	CTCACAGGCAC	0,18	1	5,56	0,49	Hs.139784	ESTs
733	TACCCATTACC	0,18	1	5,56		Hs.136981	ESTs
	TTTCTTTCCCT	0,18	1	5,56	0,49	Hs.135055	ESTs
735	CATCTGTACTG	0,18	1	5,56	0,49	Hs.132892	Homo sapiens
							protocadherin 10 (PCDH10) mRNA, partial
736	GAGCTGTTTTG	0,18	1	5,56	0,49	Hs.127476	ESTs
737	ACACGTACTAT	0,18	1	5,56	0,49	Hs.120828	Human DNA sequence
						*	from clone RP5-876B10 on
738	TTTATATTTCA	0,18	1	5,56	0,49	Hs.11958	oxidative 3 alpha
							hydroxysteroid dehydrogenase; reti
739	CACAGTTTTAA	0,18	1	5,56	0.49	Hs.118918	ESTs
	TGGTTTTACCA	0,18	1	5,56		Hs.117582	CGI-43 protein
	GTGCCTGGTAT	0,18	1	5,56		Hs.116459	hypothetical protein
710	TAGOTTA 6 : : 6					11- 44 1555	PRO2198
142	TACCTTAGAAC	0,18	1	5,56	U,49	Hs.114963	Homo Sapiens (clone

	<del></del>			<del></del> ,			Inches in
							B3B3E13) chromosome 4p16.3 DNA
743	GTGTAGGAGGT	0,18	1	5,56		Hs.113029	ribosomal protein S25
744	CCCTGGAGACA	0,18	1	5,56	0,49	Hs.111334	ferritin, light polypeptide
745	CCTGTAGCCCA	0,18	1	5,56	0,49	Hs.109370	ESTs
746	TATCCATATTA	0,18	1	5,56	0,49	Hs.107942	DKFZP564M112 protein
747	CTACTAATTGC	0,18	1	5,56		Hs.101916	Homo sapiens mRNA; cDNA DKFZp564K133 (from clone
748	GGCCTCCAAGA	0,91	5	5,49	2,25	Hs.286220	ESTs
749	GTGGCTGACAC	0,73	4	5,48	1,83	Hs.202234	ESTs, Weakly similar to ALU4_HUMAN ALU SUBFAMILY
750	CTGTACTTGTG	1,49	8	5,37	·	Hs.75678	FBJ murine osteosarcoma viral oncogene homolog B
751	ACAAAACCCCA	1,49	8	5,37	3,37	Hs.140208	ESTs
752	AGCCACCACCC	0,56	3	5,36		Hs.232045	ESTs, Moderately similar to ALU1_HUMAN ALU
753	TGCCTGTAGTT	1,32	7	5,30		Hs.246646	EST
754	GCTAACCCCTG	2,84	15	5,28	5,92	Hs.279772	brain specific protein
755	GCAAAACCCTG	16,29	86	5,28	31,14	Hs.117582	CGI-43 protein
	AAACATTAAAA	3,98	21	5,28		Hs.77443	actin, gamma 2, smooth muscle, enteric
757	GTGGCACTTGC	0,76	4	5,26	1,78	Hs.272322	Homo sapiens mRNA; cDNA DKFZp434L092 (from clone
758	GAAGCTACACC	0,76	4	5,26	1,78	Hs.107253	ESTs
759	TAAGGTAGAGG	0,38	2	5,26	0,94	Hs.99908	nuclear receptor coactivator 4
760	TGGGTCATTTG	0,38	2	5,26	0,94	Hs.98073	ESTs
761	GCTCACTGAAG	0,38	2	5,26	0,94	Hs.9568	zinc finger protein 261
762	CACCTATCAAT	0,38	2	5,26	0,94	Hs.58617	Rho-associated, coiled- coil containing protein kinas
	CTTCAATCTTA	0,38	2	5,26	0,94	Hs.58419	DKFZP586L2024 protein
	CAGTCCTCTTG	· 10,38	.2	5,26		Hs.57553	tousled-like kinase 2
	TTACCAAAGCA	0,38	2	5,26	•	Hs.30246	solute carrier family 19 (thiamine transporter), mem
	ATGAAACCCTA	0,38	2	5,26		Hs.282671	EST
	TCACTCCAGCC	0,38	2	5,26		Hs.270497	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY
	AAAGGCATCAG	0,38	2	5,26		Hs.256297	integrin, alpha 11
	GACATCTGTCC	0,38	2	5,26		Hs.25566	ESTs
	TGTCTTTTCTG	0,38	2	5,26	0,94	Hs.250882	bradykinin receptor B2
771   	TGACTGTATTA	0,38	2	5,26	0,94	Hs.198241	amine oxidase, copper containing 3 (vascular adhesio

262

772	GCAAGACCTCA	0,38	2	5,26	0.94	Hs.181592	ESTs
	AGGAAGGAAAA	0,38	2	5,26		Hs.180532	heat shock 90kD protein
' ' '	7.00,7.00,777	0,00	-	0,20	0,04	1 13. 100002	1, alpha
774	CTGCCGGAGCA	0,38	2	5,26	0.04	Hs.164779	ESTs
	TATTCCAGAAC	0,38	2	5,26		Hs.161554	hypothetical protein
113	INTICONOVIC	0,36	4	3,20	0,54	115.101334	FLJ20159
776	AGCGAAACTCC	0,38	2	5,26	0.04	Hs.106597	ESTs
	CTGGGGGTCAG	0,38	2	5,26		Hs.102664	vesicle-associated
' ' '	CIGGGGTCAG	0,36	2	5,26	0,94	ITS. 102004	membrane protein 4
778	GCTGTAATCCC	2,9	15	5,17	5.82	Hs.184019	Homo sapiens clone
' ' '		,		•,	0,02	1.0.10.10.10	23551 mRNA sequence
779	CACCTGTGGTC	3,14	16	5,10	6,11	Hs.209585	EST
780	CTCTAGAGAAA	0,59	3	5,08	1,35	Hs.97925	hypothetical protein
781	GCAAATCCTGT	0,59	3	5,08	1,35	Hs.79059	transforming growth
				·			factor, beta receptor III
							(betag
782	TATATGCTGGG	0,59	3	5,08	1,35	Hs.2969	v-ski avian sarcoma viral
			[				oncogene homolog
783	GGGAGACCCCA	0,59	3	5,08	1,35	Hs.24884	ESTs, Moderately similar
	•				,		to RNA polymerase i
			- 1	- 1			associa
784	TGGAAATGAAA	0,59	3	5,08	1,35	Hs.172928	collagen, type I, alpha 1
785	CCTGTAATCTG	0,59	3	5,08	1,35	Hs.159975	ESTs
786	TGTATTGTACA	0,59	3	5,08	1,35	Hs.118562	Link guanine nucleotide
							exchange factor II
787	ACATAGACCGA	1,38	7	5,07	2,87	Hs.173594	pigment epithelium-
							derived factor
788	GTTCCACAGAA	2,37	12	5,06		Hs.179573	collagen, type I, alpha 2
789	GTGGCAGAGAC	0,79	4	5,06	1,74	Hs.75813	polycystic kidney
							disease 1 (autosomal
							dominant)
790	AGCCGAGATCG	0,79	4	5,06	1,74	Hs.278053	EST
791	CCTGTAGTTCC	4,75	24	5,05	8,84	Hs.189242	Homo sapiens mRNA;
	•						cDNA DKFZp434A202
							(from clone
792	GTAAAACCCTG	5,77	29	5,03	10,5	Hs.281680	peroxisomal trans 2-
			1				enoyl CoA reductase;
							putative sh
	AGTCTGCTGGG	0	5	5,00	7,44	Hs.259508	ESTs
794	CTGGGCAGAGA	0	5	5,00	7,44	Hs.200735	ESTs, Moderately similar
				.			to AF105377_1 heparan
							sulfa
	TCACACAAAGG	1	5	5,00		Hs.46783	ESTs
796	CGCCTGTAATT	1	5	5,00	2,11	Hs.14333	ESTs

Tabelle 1:

Nr.	Tag_Sequence	CGAP	Rel. Expr. Haut	Quot ient	signifi c.	UniGene AccNr.	Beschreibung
798	ATTGCTCTCTG	1,61		4,97	3,18	Hs.245188	tissue inhibitor of metalloproteinase 3 (Sorsby fund
799	CCTGTGGTTCC	2,02	10	4,95	3,88	Hs.286061	ESTs
800	TGTCCACACAT	0,82	4	4,88	1,69	Hs.5897	Homo sapiens mRNA; cDNA DKFZp586P1622 (from clo
801	GACGGAGCCTT	0,82	4	4,88	1,69	Hs.37482	COPZ2 for nonclathrin coat protein zeta-COP
802	ACTGGGCAGTG	0,82	4	4,88	1,69	Hs.241257	latent transforming growth factor beta binding prote
803	CGGCACCTTAA	0,82	4	4,88	1,69	Hs.209100	DKFZP434C171 protein
804	CTGAAACAGCT	0,82	4	4,88	1,69	Hs.106469	suppressor of var1 (S.cerevisiae) 3-like 1
805	TGTTACCTGGT	0,41	2	4,88	0,9	Hs.79227	myomesin (M-protein) 2 (165kD)
806	TCTTTGCTCTT	0,41	2	4,88	0,9	Hs.44077	hypothetical protein FLJ10793
807	AAATCACCAAT	0,41	2	4,88	0,9	Hs.4082	lectin, galactoside-binding, soluble, 8 (galectin 8)
808	TGTATTGACTG	0,41	2	4,88	0,9	Hs.29280	ESTs
809	TTCACTTCAAC	0,41		4,88	0,9	Hs.250911	Homo sapiens clone 23967 unknown mRNA, partial cds
	CGTGGGGCTGC	0,41		4,88		Hs.221986	aquaporin 5
	TCACAGAGTCT	0,41	2	4,88	0,9	Hs.194149	Homo sapiens mRNA; cDNA DKFZp434M011 (from
812	TGCCACTGTGC	0,41	2	4,88	0,9	Hs.183165	ESTs, Highly similar to protein kinase [H.sapiens]
813	CTATAAAAGTG	0,41	2	4,88	0,9	Hs.154145	guanine nucleotide binding protein (G protein), alph
814	TTGGCTAGGCT	2,67	13	4,87		Hs.252259	ribosomal protein S3
815	AGGGAGCAGAG	2,26				Hs.118223	microfibrillar-associated protein 4
816	TTGGTGTGCTG	1,44	7	4,86	2,78	Hs.240399	EST
	TGGAAAGTGAA	10,31				Hs.25647	v-fos FBJ murine osteosarcoma viral oncogene homolog
	TTGATTGAGTG	0,62		4,84		Hs.9879	ESTs
	CAGGGATCTGC	0,62	3	4,84	1,31	Hs.7634	ESTs, Moderately similar to semaphorin B [M.musculus
820	ATTGCACCACC	0,62	3	4,84	1,31	Hs.44259	ESTs, Weakly similar to

							T
604	OTOOTOOOO	0.00				11 000000	ALU1_HUMAN ALU
	CTGGTGGCCAC	0,62		4,84	·	Hs.286028	Human alpha-2 collagen type VI mRNA, 3' end
	AATCATTGAGG	0,62		4,84		Hs.19150	Homo sapiens mRNA; cDNA DKFZp564A2164 (from
823	ACCCCCTTCCT	0,62	3	4,84	1,31	Hs.181392	major histocompatibility complex, class I, E
824	CCTGGCCAAAA	0,62	3	4,84		Hs.126824	EST
825	TTAACCCCTCC	6,21	30	4,83	10,46	Hs.78224	ribonuclease, RNase A family, 1 (pancreatic)
826	ACAGGCTACGG	24,47	118	4,82	39	Hs.75777	transgelin
827	AGATGAGATGA	3,57	17	4,76	6,08	Hs.285313	core promoter element binding protein
	AAAAAAAAGGC	0,21	1	4,76	0,44	Hs.90077	TG-interacting factor (TALE family homeobox)
	TACGCTAAAAC	0,21		4,76		Hs.87354	ESTs
	GTAGAAGTGTA	0,21		4,76		Hs.8705	ESTs
	TGCCCAGCAAT	0,21	1	.,,,		Hs.76297	G protein-coupled receptor kinase 6
	AGCAGGTTTGC	0,21		4,76		Hs.7434	ESTs
	AGAGAGAGCCC	0,21	1	.,		Hs.724	thyroid hormone receptor, alpha (avian erythroblasti
834	GCACTGATTAA	0,21	. 1	4,76	0,44	Hs.71741	ESTs, Highly similar to I38945 melanoma ubiquitous m
835	CAGAGACAAGC	0,21	1	4,76	0,44	Hs.71721	ESTs
836	CAGTTGTCTAG	0,21	1	4,76		Hs.58882	Microfibril-associated glycoprotein-2
837	TGTGTGTGCTA	0,21	1	4,76	0,44	Hs.55533	ESTs
838	GCCTGGACCAG	0,21		4,76		Hs.55407	Homo sapiens mRNA; cDNA DKFZp434K0621 (from
839	TGTTTAATAAA	0,21	. 1	4,76	0,44	Hs.50841	ESTs
	CATTTTCTAAT	0,21	1	4,76	0,44	Hs.48376	Homo sapiens clone HB-2 mRNA sequence
	AGCTTTCCCAA	0,21	1	4,76	0,44	Hs.45109	ESTs
	ACTCAGTAGCC	0,21	1	4,76	0,44	Hs.44197	hypothetical protein DKFZp564D0462
	TCAAATTGAAA	0,21	1	4,76	0,44	Hs.44038	pellino (Drosophila) homolog 2
	GGCCAGCCCTG	0,21		4,76		Hs.4243	ESTs
	ACATTTTGTTC	0,21		4,76		Hs.42116	ESTs
	GGCAAGCAGGC	0,21	1	4,76		Hs.41688	dual specificity phosphatase 8
847	TGCTTGGTACA	0,21	1	4,76	0,44	Hs.393	prostaglandin I2 (prostacyclin) receptor (IP)
	TAAGTCTAATT	0,21		4,76	0,44	Hs.35804	hect domain and RLD 3
849	TATTTACTTG	0,21	1	4,76	0,44	Hs.30340	hypothetical protein KIAA1165

850	CTCTCTCCCAG	0,21	1	4,76	0.44	Hs.30172	ESTs
	AGCAAATTTTC	0,21		4,76		Hs.29423	ESTs, Weakly similar to
651		0,21	'	4,70	0,44	115.25425	macrophage lectin 2
]						ŀ	[H.sapie
852	CCAAGACCTCT	0,21	1	4,76	0.44	Hs.283619	zinc finger protein 236
	GACAGGTTCTG	0,21		4,76		Hs.272023	transforming, acidic coiled-
000	SACAGOTTOTO	0,21		7,10	0,44	1 13.27 2023	coil containing protein
854	CTTCTGGAGAA	0,21	1	4,76	0.44	Hs.267263	hypothetical protein
	TGGGTTTTGTT	0,21		4,76		Hs.26243	Homo sapiens cDNA
		. 0,21		7,70	0,7-7	113.20240	FLJ11177 fis, clone
							PLACE1007402
856	CACCGCTGCAG	0,21	<u> </u>	4,76	0 44	Hs.261373	adenosine A2b receptor
		0,2.	•	'','	0,	1.0.201010	pseudogene
857	GAAACAGGAAA	0,21	1	4,76	0.44	Hs.257387	EST
	CCTTCTTGCTA	0,21		4,76		Hs.24743	hypothetical protein
		-,-	•	,,,,,	•, • •		FLJ20171
859	ATAATCTGAAG	0,21	1	4,76	0.44	Hs.2441	KIAA0022 gene product
	CTAAATATAGG	0,21		4,76		Hs.23581	leptin receptor gene-
	•	,_ ,	Ţ	.,	, ,,,,		related protein
861	AATGCTATGGT	0,21	1	4,76	0.44	Hs.23450	ESTs
	GTGAATGAAAC	0,21		4,76		Hs.223437	EST
	TCCACAAAAA	0,21		4,76		Hs.21035	KIAA1130 protein
	AGAATTGCTTA	0,21		4,76		Hs.203188	EST
<del></del>	CCTCTAATTCC	0,21	1			Hs.192949	ESTs, Moderately similar
		· i		•	•		to ALU1_HUMAN ALU
866	TTGGACAAGAA	0,21	1	4,76	0,44	Hs.189902	ESTs
867	GGCTGGGCGCG	0,21	1	4,76	0,44	Hs.188339	ESTs
868	CAACACAAAGC	0,21		4,76		Hs.185013	ESTs, Moderately similar
							to ALU7_HUMAN ALU
869	GGCCCGGCCTC	0,21	1	4,76	0,44	Hs.183994	protein phosphatase 1,
							catalytic subunit, alpha isof
870	GCTAAAAAATT	0,21	1	4,76	0,44	Hs.183760	glucose regulated protein,
		•					58kD
871	TTTTCCTTGTT	0,21	1	4,76	0,44	Hs.182937	peptidylprolyl isomerase A
						<u></u>	(cyclophilin A)
872	GCCGCCTCTGT	0,21	1	4,76	0,44	Hs.180799	ESTs, Moderately similar
							to ALU5_HUMAN ALU
	GGCTGGTTCCA	0,21	1			Hs.179943	ribosomal protein L11
	AAGGAAATGAA	0,21	1	- 11		Hs.174131	ribosomal protein L6
	AATAGATGATA	0,21	1			Hs.174104	ESTs
	TATAAGTGGAC	0,21	1	<del></del>		Hs.17301	ESTs
	GAAAACATAAT	0,21		4,76		Hs.169329	DKFZP564A043 protein
<del></del>	CTTTGTAAAAA	0,21		4,76		Hs.16578	ESTs
	AGCTGTTTAAA	0,21		4,76		Hs.164480	ESTs
880	GGGGACAGAGC	0,21	1	4,76	0,44	Hs.161554	hypothetical protein
							FLJ20159
	ATAAAGCCAAA	0,21		4,76		Hs.159471	ZAP3 protein
882	CCCCTGCATTC	0,21	1	4,76	0,44	Hs.158302	chromosome 1 open
J	1		}	] -	)	ŀ	reading frame 1

						<u> </u>	
883	GCTGAGAAGCA	0,21	1	4,76	0,44	Hs.155975	protein tyrosine
}		] ]				}	phosphatase, receptor
							type, C-assoc
884	GAATACGTTGG	0,21	1	4,76	0,44	Hs.155596	BCL2/adenovirus E1B
-005	A T T T A A C A A	0.01		1.70	0.44	11 455040	19kD-interacting protein 2
885	ATTTCTAACAA	0,21	1	4,76	0,44	Hs.155049	hypothetical protein
000	ACATOTTOTTO	0.24		4.70	0.44	110 44904	FLJ11282
800	AGATCTTCTTG	0,21	1	4,76	0,44	Hs.14894	trans-Golgi network
							protein (46, 48, 51kD isoforms)
887	AAAAACTCTAC	0,21	1	4,76	0.44	Hs.146226	ESTs
	AGGATAAACTC	0,21	<del></del> i			Hs.14427	ESTs
	CTTGTGAGGCC	0,21	1			Hs.142428	KIAA1161 protein
	GTAACAGTAAT	0,21	<u> </u>	4,76		Hs.137396	ESTs
	TAAACGAAAAT	0,21		4,76		Hs.135465	ESTs
<u> </u>	AGGGGAATGGG	0,21		4,76		Hs.134933	ESTs
	GGCCGTGCTGC	0,21		4,76		Hs.131034	ESTs
	GGAACCTATCC	0,21		4,76		Hs.128807	ESTs
-	CCAGTGCCCTC	0,21		4,76		Hs.128630	ESTs
	ATTITGCTTAA	0,21		4,76		Hs.126558	ESTs
	AGCCAAGAGCC	0,21		4,76		Hs.125877	ESTs
	GTAGACTGAAA	0,21		4,76		Hs.124165	ESTs
	TGGTATGCACC	0,21		4,76		Hs.1191	KIAA0073 protein
	TCTGTTTTGTG	0,21		4,76		Hs.118923	ESTs
	CCAATCAATGG	0,21		4,76		Hs.116674	pre-mRNA splicing factor
$\overline{}$	TTCCCCAGGCT	0,21		4,76		Hs.116296	ESTs
	CAACAGCCCCA	0,21		4,76		Hs.11455	ESTs
	CTCCTGGAATA	0,21		4,76		Hs.11367	hypothetical protein RP1-
		","	·	.,. •	٠,	,	317E23
905	TGGTGAAGAAC	0,21	1	4,76	0,44	Hs.113052	RNA cyclase homolog
	CACACCGCCCG	0,21		4,76		Hs.112015	ESTs, Moderately similar
					·		to alpha tubulin [H.sapiens
907	CCCCCCCCA	0,21	1	4,76	0,44	Hs.110953	Homo sapiens mRNA;
	ĺ ·	ĺĺ					cDNA DKFZp434A139
·							(from clone
	TTTTGTTGCTC	0,21	. 1	4,76		Hs.109641	ESTs
909	GTAAAACCCCG	5,48	26	4,74	9	Hs.258881	EST, Weakly similar to
1							ALU8_HUMAN ALU
							SUBFAMILY
	TAATAAAGAAT	0,85		4,71		Hs.80342	keratin 15
	AAAATAAACCT	0,85		4,71		Hs.74304	periplakin
	GTGGTAGGTGC_	0,85		4,71		Hs.254237	EST
913	GTGAAATTCCA	0,85	4	4,71	1,65	Hs.228168	ESTs, Weakly similar to
	00707467006	0.05		4 = 2	4.5=	11 44555	ALU1_HUMAN ALU
914	CCTCTAGTCCC	0,85	4	4,71	1,65	Hs.145501	ESTs, Weakly similar to
L-045	TOOMATOAGG	00.50	400	4 76	04.00	11- 470000	ALU1_HUMAN ALU
	TGGAAATGACC	22,53	106			Hs.172928	collagen, type I, alpha 1
916	AACCCTGGAGG	1,49	7	4,70	2,69	Hs.283927	Homo sapiens
1	l	! l		l I		}	cytomegalovirus partial

		—Т					fusion receptor
917	AGGATCACTTG	0,64	3	4,69	1 26	Hs.287355	ESTs
	CCACTGCATTG	0,64		4,69		Hs.278551	ESTs, Weakly similar to
							ALU2_HUMAN ALU
	AACTGGCTGCT	0,64		4,69		Hs.173381	dihydropyrimidinase-like 2
	CCTGTAACCCC	5,98		4,68		Hs.176541	ESTs
921	GCACTCCAGCC	6,91	32	4,63	10,69	Hs.274580	Homo sapiens mRNA; cDNA DKFZp434P1019 (from
922	GTGGTGAGCAC	1,08	5	4,63		Hs.27311	single-minded (Drosophila) homolog 2
	ATGGCAGGTGC	3,46		4,62		Hs.236479	EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY
	CCATTACACTC	1,73		4,62		Hs.8594	KIAA1191 protein
	GGTGACAGAGT	2,61		4,60		Hs.278314	hypothetical protein
926	GTGGTTTGCTG	1,96	9	4,59	3,31	Hs.157773	ESTs, Moderately similar to OPSB_HUMAN BLUE-
927	TCTGCCTATGC	0,88	4	4,55	1,6	Hs.90291	laminin, beta 2 (laminin S)
928	GCGAAAACCCC	0,88	4	4,55	1,6	Hs.272837	ESTs
929	TACCCCTTGAA	0,44	2	4,55		Hs.95834	ESTs
	TAAATGGAAGT	0,44		4,55		Hs.6942	ESTs
931	TGTGTCTGGGA	0,44		4,55		Hs.60548	hypothetical protein PRO1635
932	GCTTCAGTGGG	0,44	2	4,55	0,86	Hs.54828	ESTs
	CTTTACTGTGT	0,44		4,55		Hs.279853	HSPC018 protein
934	CACTITGTGTG	0,44		4,55		Hs.24752	spectrin SH3 domain binding protein 1
935	GGTGACAGAAC	0,44	2	4,55	0.86	Hs.234890	EST
	ACTAAGATTGA	0,44		4,55		Hs.227716	KIAA0934 protein
	CCAATGTTGTT	0,44	2	4,55	0.86	Hs.22209	ESTs
	GTGGTGTGAGC	0,44		4,55	0,86	Hs.194637	Homo sapiens mRNA; cDNA DKFZp564D113 (from clone
	TGAAGAATGTG	0,44		4,55		Hs.112557	ESTs, Moderately similar to ZN85_HUMAN ZINC
	GCCAGTGCCTG	0,44	2	4,55	0,86	Hs.106061	RD RNA-binding protein
	CCACTGCTCTC	2,64	12	4,55	4,26	Hs.23510	Kruppel-like factor 12
942	CTATTGCACTC	1,32		4,55		Hs.160483	erythrocyte membrane protein band 7.2 (stomatin)
943	ATGTACCTGAT	1,55	7	4,52	2,61	Hs.29191	epithelial membrane protein 2
944	GTGGGGGGAG	1,55	7	4,52	2.61	Hs.10700	hypothetical protein
	CACCTGTAGTC	8,23				Hs.267812	sorting nexin 4
	TCTGCACACAC	0,67		4,48		Hs.78518	natriuretic peptide receptor B/guanylate cyclase B (
947	GACAATTCTGT	0,67	3	4,48	1,22	Hs.186571	hypothetical protein FLJ10700
948	AGGGGAAGGTG	1,79	8	4,47	2,92	Hs.112540	EST

949	CAAGACGGGGG	4,31	19	4,41	6,29	Hs.106185	ral guanine nucleotide dissociation stimulator
950	TAATTTGCGTT	0,91	1	4,40	1.56	Hs.79368	epithelial membrane
330	17411100011	0,91	4	7,40	1,50	113.79300	protein 1
951	AGAATCGTTTG	0,91	4	4,40	1,56	Hs.136299	ESTs ·
952	AACAGTCAAAA	3,87		4,39		Hs.26557	plakophilin 3
	GCAAAACTCTG	2,29		4,37		Hs.278746	ESTs, Moderately similar to ALU8_HUMAN ALU
954	GCCAGCCAGTG	2,29	10	4,37	3.49	Hs.149098	smoothelin
	GCGAAATCCCA	1,38		4,35		Hs.268728	ESTs
	TCTGTGGTCCC	1,38		4,35		Hs.227894	ESTs, Weakly similar to ALU1_HUMAN ALU
957	CCTATAATTCC	1,38	6	4,35	2,21	Hs.135491	ESTs
958	GCAATAAAATA	0,23	1	4,35	0,4	Hs.99621	ESTs
959	TGGAATCCAGG	0,23	1	4,35	0,4	Hs.98135	ESTs
	CTTTTGTCAGC	0,23		4,35		Hs.90858	Homo sapiens clone
					•	1	25023 mRNA sequence
961	TTATTGTATTG	0,23	1	4,35	0,4	Hs.89474	ADP-ribosylation factor 6
962	ATGGAGGTATG	0,23		4,35		Hs.8944	procollagen C-
		. ,		,			endopeptidase enhancer 2
963	TCGTGTTTTCG	0,23	1	4,35	0,4	Hs.87595	translocase of inner
		.,		-,			mitochondrial membrane
					_		22 (yeas
964	TTGTGGCCCCA	0,23	1	4,35	0,4	Hs.84630	ESTs
	TITGTTTGTTT	0,23		4,35		Hs.8355	ESTs
	GTCACGAACAT	0,23		4,35		Hs.82933	protein x 013
	GTTAAGGTAAA	0,23		4,35		Hs.79241	B-cell CLL/lymphoma 2
	AAGAAGAAAAG	0,23		4,35		Hs.78293	ESTs
	TAGCCAGTTAA	0,23		4,35		Hs.74101	spleen tyrosine kinase
	CTGCCCGGGGC	0,23		4,35		Hs.74097	mercaptopyruvate
			· , _				sulfurtransferase
	AACGGTGTTTG	0,23		4,35		Hs.71371	ESTs
	CATAAACGGGC	0,23		4,35		Hs.69954	laminin, gamma 3
	TCCTTAGATTA	0,23		4,35		Hs.69743	GM2 ganglioside activator protein
	CTGGCATAGAA	0,23		4,35		Hs.61272	ESTs
	ACTGCCCCTGA	0,23	1	4,35	0,4	Hs.59729	semaphorin sem2
	TCTGCTGCCTG	0,23		4,35		Hs.58006	ESTs
977	GAAGATGAATA	0,23	1	4,35	0,4	Hs.54982	ESTs, Weakly similar to ALU4_HUMAN ALU
978	GACCAAAGAAG	0,23	1	4,35	0,4	Hs.48948	ESTs
	GAATGAATGCA	0,23		4,35		Hs.48604	DKFZP434B168 protein
980	GCACAACTAAA	0,23		4,35		Hs.47587	ESTs
	GAATTTTACAC	0,23		4,35		Hs.47522	ESTs
	AAGGGGCGCG	0,23		4,35		Hs.456	leukotriene C4 synthase
	AGGGACTTTAT	0,23		4,35		Hs.43148	ESTs
	TATTCAGAACC	0,23		4,35		Hs.40289	ESTs
	ACAACTGGAAT	0,23		4,35		Hs.37372	Homo sapiens DNA
		-,=-	Ī	"			binding peptide mRNA,

							partial cds
	TAATAAAATGC	0,23	1	4,35		Hs.29008	ESTs
987	TTATGCTTGTA	0,23	1	4,35	0,4	Hs.284153	Fanconi anemia,
							complementation group A
988	TTTTGAAGATA	0,23	1	4,35	0,4	Hs.283322	hypothetical protein
989	GCAAATCAGAT	0,23	1	4,35	0,4	Hs.279477	ESTs
990	GTAAAACCCCT	0,23	1	4,35	0,4	Hs.277896	EST
991	GGCCAGGCGTG	0,23	1	4,35	0,4	Hs.276994	EST
	AATGTTAGAGC	0,23		4,35		Hs.270331	ESTs
993	TACCTATAGTC	0,23				Hs.269838	ESTs .
	TGTGAGAAAGT	0,23	1	4,35		Hs.241493	natural killer-tumor
		,					recognition sequence
995	AATTGTGCATT	0,23	1	4,35	0.4	Hs.240443	chondroitin 4-
		5,	·	.,	-,		sulfotransferase
996	GTGGTGCGCAT	0,23	1	4,35	0.4	Hs.236505	EST, Weakly similar to
		5,25	•	.,	٠,,		ALU6_HUMAN ALU
		i					SUBFAMILY
997	GTTTATAATTA	0,23	1	4,35	0.4	Hs.231966	ESTs
	CTGGCACCCTG	0,23	1			Hs.212716	EST
	AAAAATGGTGG	0,23	1	4,35		Hs.204930	EST
	GAGCAGGCAAA	0,23	1	4,35		Hs.200333	apolipoprotein B48
.000		0,20	•	1,00	0, 1	1.0.20000	receptor
1001	CCAAAAAGTG	0,23	1	4,35	0.4	Hs.184242	sterol-C5-desaturase
,00,		0,20		1,00	0, 1	110.10-12-12	(fungal ERG3, delta-5-
							desaturas
1002	CCAGAGGAATG	0,23	1	4,35	0.4	Hs.180414	heat shock 70kd protein
.002		0,20	•	1,00	0,4	113.100414	10 (HSC71)
1003	CCACAAAAAA	0,23	1	4,35	0.4	Hs.179091	EST
	GCTTACCTGCT	0,23	<del></del>	4,35		Hs.174031	cytochrome c oxidase
1001	00177.001001	0,20		7,00	.0,-1	113.174001	subunit VIb
1005	TATTAGACACC	0,23	1	4,35	0.4	Hs.17258	ESTs
	GAATGTTGACA	0,23	1			Hs.16959	ESTs
	TGAGGGGTGGG	0,23	1			Hs.166293	EST
	TATATAAGTAC	0,23	<del></del>	4,35		Hs.166011	catenin (cadherin-
1000		0,23		4,55	0,4	115.100011	associated protein), delta
							1
1009	TAATAATACAA	0,23	1	4,35	0.4	Hs.16349	KIAA0431 protein
	TGGGAGGCTGA	0,23	1	4,35		Hs.161554	hypothetical protein
1010	IGGGAGGCIGA	0,23	,	4,55	0,4	ITS. 10 1334	FLJ20159
1011	AGCTCCTTAAG	0.22	1	4,35	0.4	Un 150500	
	CTTTTTGTGGT	0,23	1			Hs.159509	alpha-2-plasmin inhibitor
1012		0,23	I	4,35	0,4	Hs.153106	Homo sapiens clone
1012	CTGTTTTTGAA	0.22	1	1 25	0.4	Un 152720	23728 mRNA sequence
	GCAGACTATCC	0,23				Hs.152720	M-phase phosphoprotein 6
		0,23		.,		Hs.151696	DKFZP727G051 protein
	GTTCCCTGGTG	0,23		4,35		Hs.146090	ESTS
1016	CCTACAGTCCC	0,23	1	4,35	υ,4	Hs.144874	Homo sapiens mRNA;
						j	cDNA DKFZp761C0524
4047	CAATOOTOOTO	- 0.00		4.05		11- 400 40=	(from
1017	CAATCCTCCTG	0,23	1	4,35	<sub> </sub> 0,4	Hs.138407	ESTs

		,					
	AACAAGTAATA	0,23		4,35		Hs.134350	ESTs
	TCCAGTACAGA	0,23		4,35		Hs.12969	hypothetical protein
1020	GGAGGCTGGGG	0,23	1	4,35	0,4	Hs.127452	ESTs
1021	ACTCTGCTCGG	0,23	1	4,35	0,4	Hs.126900	ESTs
1022	TTCTGTGAGTG	0,23		4,35		Hs.122559	ESTs
	TCCTGTGATTT	0,23		4,35		Hs.12253	ESTs
	AAAGCACAAGT	0,23		4,35		Hs.111758	keratin 6B
	TTTTTGAAAAA	0,23	1	4,35		Hs.109646	NADH dehydrogenase
		,	·	',"	0, .	1.0.   0.00 1.0	(ubiquinone) 1 beta
				[ [		ĺ	subcomplex, 6
1026	AAGTCCTGGCC	0,23	1	4,35	0.4	Hs.109314	ESTs, Weakly similar to
		J,	•	.,	٥, .		2202255A AT motif-
		1					binding fa
1027	TGTTACTGGAT	0,23	1	4,35	.0.4	Hs.100861	ESTs, Weakly similar to
		0,20	·	',00	٠, ١	1.10.100001	spastin protein [H.sapiens]
1028	CCTGTAATCTT	1,85	8	4,32	2 84	Hs.120882	ESTs, Moderately similar
		1,00	J	1,02	2,0	1.0.120002	to ALU1_HUMAN ALU
1029	GTGGCGGCAC	19,81	85	4,29	25 22	Hs.283044	hypothetical protein
.020	0.0000000000000000000000000000000000000	10,01		1,20	20,22	7.3.2000-7-7	PRO2859
1030	CCACTTGCACT	0,7	3	4,29	1 18	Hs.220962	EST, Weakly similar to
1000	00/10/100/10/	٠,,	•	7,20	1,10	113.220302	ALU1_HUMAN ALU
[				ĺĺ			SUBFAMILY
1031	TTCCATACCCC	0,7	3	4,29	1 18	Hs.180398	LIM domain-containing
	1.100,117,0000	0,,,	J	7,20	1,10	113.100000	preferred translocation
			•				partne
1032	ATTGCATCACT	1,17	5	4,27	1 85	Hs.209111	EST
	GAGTTAAAAAA	1,17	5	4,27		Hs.180255	major histocompatibility
		.,	Ĭ	.,	.,00	10.100200	complex, class II, DR beta
1034	GGGCCCTGGCC	0,94	4	4,26	1.53	Hs.25895	ESTs, Weakly similar to
		-,-	Ì	',==	.,		PI-3 kinase [H.sapiens]
1035	CAGATGGAGGC	0,94	4	4,26	1.53	Hs.127273	hypothetical protein
		١,٠٠١	j	',	.,		FLJ10044
1036	CCTCTCCCACA						
	ICCICICCCACA I	0.471	2	4.26	0.82	Hs.99197	
1 1037		0,47	2	4,26 4,26		Hs.99197 Hs.96875	ESTs
	GCCAGGGGGTA	0,47	2	4,26	0,82	Hs.96875	ESTs ESTs
			2	4,26 4,26 4,26	0,82		ESTs ESTs CCAAT/enhancer binding
1038	GCCAGGGGGTA CTCAGTCTTTT	0,47 0,47	2	4,26 4,26	0,82 0,82	Hs.96875 Hs.76722	ESTs ESTs CCAAT/enhancer binding protein (C/EBP), delta
1038	GCCAGGGGGTA	0,47	2	4,26	0,82 0,82	Hs.96875	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to
1038	GCCAGGGGGTA CTCAGTCTTTT	0,47 0,47	2	4,26 4,26	0,82 0,82	Hs.96875 Hs.76722	ESTs ESTs CCAAT/enhancer binding protein (C/EBP), delta ESTs, Highly similar to AF151903_1 CGI-145
1038 1039	GCCAGGGGGTA CTCAGTCTTTT GATGTATTCTA	0,47 0,47 0,47	2 2	4,26 4,26 4,26	0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844	ESTs ESTs CCAAT/enhancer binding protein (C/EBP), delta ESTs, Highly similar to AF151903_1 CGI-145 protein [
1038 1039	GCCAGGGGGTA CTCAGTCTTTT	0,47 0,47	2 2	4,26 4,26	0,82 0,82 0,82	Hs.96875 Hs.76722	ESTs ESTs CCAAT/enhancer binding protein (C/EBP), delta ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator
1038 1039 1040	GCCAGGGGGTA CTCAGTCTTTT GATGTATTCTA CCCTTCTGTAA	0,47 0,47 0,47	2 2	4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716	ESTs  ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s
1038 1039 1040 1041	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA	0,47 0,47 0,47 0,47	2 2 2	4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs
1038 1039 1040 1041	GCCAGGGGGTA CTCAGTCTTTT GATGTATTCTA CCCTTCTGTAA	0,47 0,47 0,47	2 2 2	4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs  ESTs, Moderately similar
1038 1039 1040 1041 1042	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA GGCCTGTAATC	0,47 0,47 0,47 0,47 0,47	2 2 2 2	4,26 4,26 4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802 Hs.267400	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs  ESTs, Moderately similar to ALU7_HUMAN ALU
1038 1039 1040 1041 1042	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA	0,47 0,47 0,47 0,47	2 2 2 2	4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs)  ESTs, Moderately similar to ALU7_HUMAN ALU Homo sapiens mRNA;
1038 1039 1040 1041 1042	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA GGCCTGTAATC	0,47 0,47 0,47 0,47 0,47	2 2 2 2	4,26 4,26 4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802 Hs.267400	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs  ESTs, Moderately similar to ALU7_HUMAN ALU Homo sapiens mRNA; cDNA DKFZp434D0215
1038 1039 1040 1041 1042 1043	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA GGCCTGTAATC  AGGTATATATC	0,47 0,47 0,47 0,47 0,47 0,47	2 2 2 2	4,26 4,26 4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802 Hs.267400 Hs.24715	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs  ESTs, Moderately similar to ALU7_HUMAN ALU Homo sapiens mRNA; cDNA DKFZp434D0215 (from
1038 1039 1040 1041 1042 1043	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA GGCCTGTAATC  AGGTATATATC  TTCTGAAAGGA	0,47 0,47 0,47 0,47 0,47 0,47	2 2 2 2 2 2	4,26 4,26 4,26 4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802 Hs.267400 Hs.24715	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs  ESTs, Moderately similar to ALU7_HUMAN ALU  Homo sapiens mRNA; cDNA DKFZP434D0215 (from DKFZP586F1019 protein
1038 1039 1040 1041 1042 1043 1044 1045	GCCAGGGGGTA CTCAGTCTTTT  GATGTATTCTA  CCCTTCTGTAA  TGGAACTGTCA GGCCTGTAATC  AGGTATATATC	0,47 0,47 0,47 0,47 0,47 0,47	2 2 2 2 2 2 2 2	4,26 4,26 4,26 4,26 4,26 4,26	0,82 0,82 0,82 0,82 0,82 0,82 0,82 0,82	Hs.96875 Hs.76722 Hs.75844 Hs.75716 Hs.285802 Hs.267400 Hs.24715 Hs.227209 Hs.154919	ESTs  CCAAT/enhancer binding protein (C/EBP), delta  ESTs, Highly similar to AF151903_1 CGI-145 protein [ plasminogen activator inhibitor, type II (arginine-s ESTs  ESTs, Moderately similar to ALU7_HUMAN ALU Homo sapiens mRNA; cDNA DKFZp434D0215 (from

	<del></del>		т			T	
10:=	0.474.477070		<u> </u>	1.55			phosphatase 2
	CATAATTTCTC	0,47		4,26		Hs.104660	elF-5A2 protein
	TGGGACGTGAG	1,67	7	4,19	2,45	Hs.3796	EphB6
	GCGAAATCCCG	2,4	· .	4,17		Hs.194251	ESTs, Weakly similar to ALU2_HUMAN ALU
	ACCAAAAACCA	28,92				Hs.172928	collagen, type I, alpha 1
	ÄACCCAGGAGG	30,47		4,14		Hs.161554	hypothetical protein FLJ20159
1052	TCTCTGTGTAG	0,97	4	4,12		Hs.79187	coxsackie virus and adenovirus receptor
	CGCAGTAGGGG	0,97		4,12		Hs.17411	KIAA0699 protein
1054	CGAGAGGGAGA	0,97	4	4,12	1,49	Hs.158159	FAT tumor suppressor (Drosophila) homolog 2
1055	GTGGCATCTGC	0,97	4	4,12	1,49	Hs.1244	CD9 antigen (p24)
1056	CTAACGGGGCG	1,7	7	4,12	2,42	Hs.102171	immunoglobulin superfamily containing leucine-rich r
1057	GCAAAACCCCT	1,46	6	4,11	2,1	Hs.75238	chromatin assembly factor 1, subunit B (p60)
1058	ACTGCTTTACT	1,46	6	4,11	2,1	Hs.72157	DKFZP564I1922 protein
1059	CCCCAGGCTGC	0,73	3	4,11		Hs.9645	ESTs
1060	TAAAATGTTTA	0,73	3	4,11	1,15	Hs.94109	ESTs
	CCTACTGCACT	0,73	3	4,11		Hs.225641	ESTs, Moderately similar to KIAA0680 protein [H.sapi
1062	GTGGCTCATTC	0,73	3	4,11	1,15	Hs.116577	prostate differentiation factor
1063	AAGCACAAAAA	2,93	12	4,10	3,87	Hs.9963	TYRO protein tyrosine kinase binding protein
1064	GTGGCGGGCGC	22,97	94	4,09	26,43	Hs.129710	malignancy-associated protein
<u> </u>	AGAACCTTAAA	3,43	14	4,08	4,43	Hs.181244	major histocompatibility complex, class I, A
1066	CCTGAAATCCC	1,96		4,08	2,69	Hs.182124	ESTs
1067	GCGAAACCCAG	3,22	13	4,04	4,1	Hs.142442	HP1-BP74
	ATGTAGGTGCC	2,23		4,04		Hs.173717	phosphatidic acid phosphatase type 2B
	TTAAATAGCAC	2,49		4,02		Hs.172928	collagen, type I, alpha 1
	TTTATTTCCA	0		4,00		Hs.93780	ESTs, Weakly similar to ALUB_HUMAN IIII ALU CLASS
	AATTACAGCCA	0		4,00		Hs.74471	gap junction protein, alpha 1, 43kD (connexin 43)
	CCTTACCTAAG	0		4,00		Hs.240217	dopachrome tautomerase (dopachrome delta- isomerase,
	CTCCCTGAACG	0		4,00		Hs.11006	ESTs
	AACACGAATGA	1		4,00		Hs.259855	ESTs
1075	GTGGCAAGCAC	1	4	4,00	1,45	Hs.138860	Rho GTPase activating protein 1

1076	TTCACCATCCT	1	4	4,00	1,45	Hs.101395	ESTs, Weakly similar to tetraspan NET-4 [H.sapiens]
1077	AGAGGGAGTGA	0,5	2	4,00	0,78	Hs.85201	C-type (calcium dependent, carbohydrate-recognition
1078	AGTCCTTGAAA	0,5	2	4,00	0,78	Hs.81665	v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogen
1079	CCCAGCCTAAA	0,5	_	4,00		Hs.47986	Homo sapiens mRNA; cDNA DKFZp586H051 (from clone
1080	TTTAACTGACA	0,5	2	4,00		Hs.24880	ESTs
1081	CCTTGTAATCC	0,5	2	4,00	0,78	Hs.197054	EST
	GAAGGCTTATC	0,5		4,00		Hs.172674	nuclear factor of activated T-cells, cytoplasmic 3
	TAGCAAAGATT	0,5		4,00		Hs.166172	aryl hydrocarbon receptor nuclear translocator
. 4	CACATCTGTAA	0,5		4,00		Hs.16533	myosin phosphatase, target subunit 1
	TTCAGTAATAA	0,5		4,00		Hs.13479	hypothetical protein FLJ20847
	GGTGAAACCCT	0,5	2	4,00		Hs.117582	CGI-43 protein
	CTTCTGCCTCA	0,5	2	4,00	0,78	Hs.115896	ESTs
1088	GAGAGGTGATT	0,5	2	4,00	0,78	Hs.114062	protein tyrosine phosphatase-like (proline instead o
1089	ATGTATTTTA	0,5	2	4,00	0,78	Hs.108396	ALR-like protein
1090	CTATAGGAGAC	1,26	5	3,97		Hs.8966	integral membrane protein
	TCCGTGTATAA	1,26		3,97	·	Hs.3321	ESTs, Highly similar to IRX3_MOUSE IROQUOIS-
	GCAAAACCCCA	24,76				Hs.129708	tumor necrosis factor (ligand) superfamily, member 1
	TTCCATAGCCT	1,52		3,95		Hs.8546	Notch (Drosophila) homolog 3
	CTGTGAAATGC	0,76		3,95		Hs.23618	hypothetical protein FLJ10704
	AAAGAACATAG	0,76		3,95		Hs.104558	ESTs
	CACCTGTAATC	9,41		3,93		Hs.275819	EST
	GGCAACAAGAG	2,29	•	3,93		Hs.205739	ESTs, Weakly similar to ALU7_HUMAN ALU
	GTGGCGGGTGC	18,52		3,89		Hs.277015	EST
	ACCTTCAAAAA	1,03	4	3,88		Hs.28444	hypothetical protein FLJ10567
	ACATCTGGCTT	1,03		3,88		Hs.194035	KIAA0737 gene product
1101	GTACGTATTCT	1,55	6	3,87	2	Hs.76325	immunoglobulin J polypeptide, linker protein

							for imm
1102	ATCCGCCTGCC	1,55		3,87		Hs.167956	ESTs, Weakly similar to
1102	AICCGCCIGCC	1,55	U	3,01	2		KIAA0309 [H.sapiens]
4400	ACCCACCTOAC	0.04	- 04	2.00	C 04	LID 409054	1
	ACCCACGTCAG	6,21		3,86		Hs.198951	jun B proto-oncogene
	TAATCCCAGCT	3,63		3,86		Hs.238384	EST
	ATTGCACCACT	11,93		3,86		Hs.117582	CGI-43 protein
	AGGACCAAGGA	0,26	1	3,85		Hs.99539	ESTs, Weakly similar to ALU7_HUMAN ALU
1107	TAAGCTACTAA	0,26	1	3,85	0,37	Hs.97469	ESTs, Weakly similar to I49698 alpha-1,3-galactosylt
1108	CTCCATTGTCT	0,26	1	3,85	0,37	Hs.93005	slug (chicken homolog), zinc finger protein
1109	TGACATTAAAC	0,26	1	3,85	0,37	Hs.87432	ESTs
	GGATTCAAGAG	0,26	1	3,85	0,37	Hs.86947	a disintegrin and metalloproteinase domain 8
1111	ATGTTATCATA	0,26	1	3,85		Hs.8325	mitogen-activated protein kinase 9
1112	TACTCTGTTGA	0,26	1	3,85		Hs.82587	phospholipase D1, phophatidylcholine-specific
1113	GGAAAAGAAAA	0,26	1			Hs.82141	Human clone 23612 mRNA sequence
1114	TATACGTTATG	0,26	1	3,85	0,37	Hs.78894	KIAA0161 gene product
1115	CACTTGGTGAT	0,26	1	3,85	0,37	Hs.77348	hydroxyprostaglandin dehydrogenase 15-(NAD)
1116	ATGAATGTAAA	0,26	1	3,85	0,37	Hs.76853	Homo sapiens mRNA full length insert cDNA clone EURO
1117	TGTTTCATAA	0,26	1	3,85	0,37	Hs.75703	small inducible cytokine A4 (homologous to mouse Mip
1118	AGTTTACGATT	0,26	1	3,85	0,37	Hs.74649	cytochrome c oxidase subunit VIc
1119	GTGGTTCATTC	0,26	1	3,85	,	Hs.6853	carbohydrate (N- acetylglucosamine 6-O) sulfotransfer
1120	CAACAAAAGCA	0,26	1	3,85		Hs.66450	ESTs
1121	CATTTTTTGCG	0,26	1	3,85	0,37	Hs.59525	ESTs
1122	CTGCTAAACTA	0,26	1	3,85		Hs.46826	ESTs .
	ACCCTGAATGG	0,26	1			Hs.43086	ESTs
	TTGTAACAAAA	0,26	1	3,85		Hs.40154	jumonji (mouse) homolog
	TGCTGTTCATA	0,26		3,85		Hs.37958	ESTs
	TATGTGGGTTA	0,26	1			Hs.34359	ESTs
	CAGCAATTATA	0,26	1			Hs.32309	inositol polyphosphate-1- phosphatase
1128	CTCCATTGCCA	0,26	1	3,85	0.37	Hs.31869	ESTs
	GGGTGGGTCAC	0,26		3,85		Hs.31500	ESTs
	ATTCCACCACT	0,26		3,85		Hs.2934	ribonucleotide reductase
	للربر	þ -, <b></b> (	•		","		M1 polypeptide

		,					.,
	TCTTACTCAGA	0,26	1	3,85		Hs.285081	ESTs
1132	AATAAAAAATA	0,26	1	3,85	0,37	Hs.284275	Homo sapiens PAK2
							mRNA, complete cds
1133	GAGACAGTGAC	0,26	1	3,85	0,37	Hs.284146	hypothetical protein
							DKFZp762N0610
	CCTGGGCAACA .	0,26	1	3,85	0,37	Hs.273683	EST
1135	CCTTTGCACTC	0,26	1	3,85	0,37	Hs.265124	ESTs
1136	TGCAGACAGGG	0,26	1	3,85	0,37	Hs.264363	hypothetical protein
							FLJ10110
1137	TTGGCCCAGTC	0,26	1	3,85	0,37	Hs.25951	Rho guanine nucleotide
							exchange factor (GEF) 3
1138	TCTTTAAAGTA	0,26	1	3,85	0,37	Hs.25155	guanine nucleotide
Ì	·	1					regulatory protein
							(oncogene)
	ATATTGGTGGT	0,26		3,85		Hs.250692	hepatic leukemia factor
	GTGGCGAATGC	0,26	1	3,85		Hs.230479	EST
1141	TATTAACATTC	0,26	1	3,85	0,37	Hs.226573	inhibitor of kappa light
							polypeptide gene
	·	<u> </u>		·			enhancer i
1142	TCTCCATTCCT	0,26	1	3,85	0,37	Hs.226573	inhibitor of kappa light
							polypeptide gene
		17.					enhancer i
	TATATAGAATG	0,26	1	3,85		Hs.22581	ESTs
1144	GCGAGATCCTG	0,26	1	3,85	0,37	Hs.22529	ESTs, Weakly similar to
							ALU1_HUMAN ALU
1145	CACTCCAACCT	0,26	1	3,85	0,37	Hs.205899	ESTs, Weakly similar to
							ALU5_HUMAN ALU
1146	CGATGTTAAAA	0,26	1	3,85	0,37	Hs.20072	myosin regulatory light
							chain interacting protein
1147	CCTTCTGAATA	0,26	, 1	3,85	0,37	Hs.194660	ceroid-lipofuscinosis,
			٠.				neuronal 3, juvenile
							(Batten,
	TGAGGATCCAG	0,26	1	3,85		Hs.191621	ESTs
1149	ATTGTACAACA	0,26	. 1	3,85	0,37	Hs.184326	CDC10 (cell division cycle
							10, S. cerevisiae, homolo
1150	AGCCTATTAAA	0,26	1	3,85	0,37	Hs.183593	zinc finger protein 24
							(KOX 17)
	GCCCCTGCGCC	0,26		3,85		Hs.183202	ESTs
	GTGAATGCACT	0,26		3,85		Hs.176065	ESTs
	TTCATTAAGAA	0,26		3,85		Hs.17411	KIAA0699 protein
	ACCGAGGTGCA	0,26		3,85		Hs.171882	ESTs
	TTGGTATTGCA	0,26	1	3,85	0,37	Hs.163541	ESTs
1156	ATGTATTTTCA	0,26	1	3,85	0,37	Hs.161554	hypothetical protein
							FLJ20159
1157	GTTGAATTGCA	0,26	1	3,85	0,37	Hs.156828	Homo sapiens cDNA
							FLJ10522 fis, clone
1158	TCTGTCATCCC	0,26	1	3,85	0,37	Hs.150158	ESTs
	GTGGTGCAAAC	0,26	1			Hs.149852	EST, Weakly similar to
							ALU1_HUMAN ALU
							SUBFAMILY
						•	•

1160	GGCTTTGGAAT	0,26	1	3,85		Hs.146481	ESTs
1161	TTTGCTTTTGA	0,26	1	3,85	0,37	Hs.144504	hypothetical protein FLJ10624
1162	AAAGCATTAGA	0,26	1	3,85	0,37	Hs.14155	KIAA0653 protein
1163	AAACAACCCCA	0,26	1	3,85		Hs.1369	decay accelerating factor for complement (CD55, Crom
1164	CTCCTTGTCCC	0,26	1	3,85	0,37	Hs.135530	EST
1165	GTGCTGCACAC	0,26	1	3,85	0,37	Hs.133272	ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS
1166	TTTAAACTTGG	0,26	1	3,85	0,37	Hs.12431	ESTs
1167	CGGCCCAGGTT	0,26	1	3,85	0,37	Hs.122823	thousand and one amino acid protein kinase
1168	CTGACTGATGG	0,26	1	3,85	0,37	Hs.121509	collagen, type XI, alpha 2
	ATAGATACACA	0,26	1			Hs.120850	ESTs
	TGGCAGTAGTG	0,26	1	3,85		Hs.120644	ESTs
1171	AAATCCTTCTA	0,26	1	3,85	0,37	Hs.119301	S100 calcium-binding protein A10 (annexin II ligand,
	TCAAACTTTGT	0,26	1	3,85		Hs.117582	CGI-43 protein
	TACACCTGGAA	0,26		3,85		Hs.114624	ESTs
	CAAGGATTTT	0,26	1			Hs.111323	Protein inhibitor of activated STAT X
	CTAGTATAAAA	0,26	1	0,00		Hs.106650	hypothetical protein FLJ20533
	AATATAAAAA	0,26	1			Hs.103548	ESTs, Weakly similar to ALU1_HUMAN ALU
	CAAATATGGTT	0,26	- 1	-,		Hs.10351	KIAA0308 protein
*	CAAGAACAGGG	0,26	1	3,85	·	Hs.102135	signal sequence receptor, delta (translocon-associat
1179	TCACCGGTCAG	15,94	61	3,83	16,2	Hs.80562	gelsolin (amyloidosis, Finnish type)
	GTGAAAACCCC	1,05		3,81	·	Hs.277213	EST, Weakly similar to ALU7_HUMAN ALU SUBFAMILY
1181	GTGGTGGCAC	15,76	60	3,81	15,85	Hs.77510	isovaleryl Coenzyme A dehydrogenase
	GTGGTGGGTGC	15,79	60	3,80	15,81	Hs.136509	EST
1183	CCGTTGCACTC	1,58		3,80		Hs.278329	ESTs
L	CATCACGGATC	0,79	3	3,80	1,08	Hs.82112	interleukin 1 receptor, type
	GTATGTACAGG	0,79	3	3,80	1,08	Hs.164255	ESTs, Moderately similar to ALU2_HUMAN ALU
	CCCTTTATATC	0,79		3,80		Hs.13766	ESTs
	GCTCGTGGTCA	0,79	3	3,80		Hs.119475	cold inducible RNA- binding protein
L	CCCATCTAGCT	0,79	3	3,80	1,08	Hs.106070	cyclin-dependent kinase inhibitor 1C (p57, Kip2)
1189	TCTCAAAAAAA	2,11	8	3,79	2,52	Hs.194841	ESTs, Moderately similar

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					L		to ALU8_HUMAN ALU
1190	CCCCTGGCTGG	1,32	5	3,79	1,68	Hs.920	modulator recognition factor I
1191	GGTTATTTAGT	1,32	5	3,79	1,68	Hs.8110	adducin 3 (gamma)
1192	GCAAAACCTCA	1,85	7	3,78		Hs.156596	ESTs, Weakly similar to ALU7_HUMAN ALU
1193	AGTTGTTTGGT	0,53	2	3,77	0,75	Hs.96418	ESTs
1194	TCATAGCCTTG	0,53	2	3,77		Hs.78846	heat shock 27kD protein 2
1195	AGGACTGGACT	0,53	2	3,77	0,75	Hs.75258	H2A histone family, member Y
	TAAACCTAGGA	0,53	2		0,75	Hs.56186	EGF-like-domain, multiple
	GTGGCTCACTT	0,53		3,77		Hs.285616	ESTs
1198	TCATTTGGTGT	0,53		3,77		Hs.285439	ESTs
1199	GCCTTGGCAGT	0,53	2	3,77	0,75	Hs.25351	iroquois-class homeodomain protein
1200	CCCTTGTTCTT	0,53		3,77	0,75	Hs.250723	FK506 binding protein 12- rapamycin associated protei
1201	GAACAGTATGA	0,53	2	3,77	0,75	Hs.189762	ESTs
1202	ATGGCAGGCGG	0,53	2	3,77		Hs.161554	hypothetical protein FLJ20159
	ACACAGCAAGA	36,07	136	·		Hs.80562	gelsolin (amyloidosis, Finnish type)
	GATCAGGCCAG	7,97		3,76		Hs.119571	collagen, type III, alpha 1 (Ehlers-Danlos syndrome
	GTGAAACTCTG	8,58		3,73		Hs.188853	Homo sapiens cDNA FLJ10150 fis, clone
	GGCCTGCAGGA	1,61	6	3,73	1,93	Hs.71869	apoptosis-associated speck-like protein containing a
1207	AAATCAATACA	1,88	7	3,72	2,21	Hs.94953	ESTs, Highly similar to . C1QC_HUMAN COMPLEMENT
1208	ATTGTACCACT	1,88	7	3,72	2,21	Hs.7099	hypothetical protein FLJ20265
	CGCCTGTAGTC	7,79	29	3,72		Hs.60088	ESTs
	GCAAAACCCAG	1,08		3,70	1,35	Hs.210610	ESTs, Moderately similar to ALU6_HUMAN ALU
	CTTTGATGCGG	1,08	4	3,70	1,35	Hs.183601	regulator of G-protein signalling 16
	GGCCCTAGGCA	8,94	33	3,69	8,73	Hs.78909	butyrate response factor 2 (EGF-response factor 2)
	CCTGGCTAATT	7,88		3,68		Hs.25661	ESTs, Weakly similar to ALUF_HUMAN !!!! ALU CLASS
	CTTCCTGGCCT	2,72		3,68		Hs.83623	nuclear receptor subfamily 1, group I, member 3
1215	GCGGGGTGGAG	10,08	37	3,67	9,65	Hs.85155	butyrate response factor 1 (EGF-response factor 1)

1216	GTGGCAGGCGC	23,18	85	3 67	21 18	Hs.48604	DKFZP434B168 protein
	AGCCCAGGAGG	2,46		3,66	2 60	Hs.136340	ESTs, Weakly similar to
'4''	ACCOCAGGAGG	2,40	9	3,00	2,09	11.15.150540	unnamed protein product
1		]			i		[H.s
1218	ATAGTGCCACT	1,64	6	3,66	10	Hs.246717	ESTs, Weakly similar to
1210	AMOTOGOAGT	',07	·	0,00	1,0	113.240717	ALU7 HUMAN ALU
1219	CATTTGTAAAA	0,82	3	3,66	1.05	Hs.84429	KIAA0971 protein
	CGTACAGCCCC	0,82	3	3,66	1.05	Hs.32580	KIAA1448 protein
	GGGCTACGTCC	0,82	3	3,66	1.05	Hs.123107	kallikrein 1,
		5,52	_	","	.,	1.01.20.01	renal/pancreas/salivary
1222	ATCACACCACT	7,41	27	3,64	7.14	Hs.234786	KIAA0707 protein
	CACTCCAGCCT	3,57		3,64		Hs.193451	ESTs, Weakly similar to
		5,51		', '	0,00		ALU7_HUMAN ALU
1224	CTTGTAATCCC	12,66	46	3,63	11.69	Hs.183253	ESTs, Weakly similar to
'		-, -, -		-, -	,		ALU1_HUMAN ALU
1225	GTGAAACCTCA	12,95	47	3,63	11,91	Hs.153029	ESTs
	ATCTCAGCTCA	3,31		3,63		Hs.246192	ESTs, Weakly similar to
					•		RMS1_HUMAN
L							REGULATOR
1227	TATGTGCTGTA	1,38	5	3,62	1,61	Hs.20084	retinoid X receptor, alpha
1228	ACTCGAATATC	1,11	4	3,60	1,32	Hs.95655	secreted and
	·				_		transmembrane 1
1229	GAGTCCCTGGT	1,11	4	3,60	1,32	Hs.68398	period (Drosophila)
:					٠,		homolog 1
1230	TGCAATATGCC	1,67	6	3,59	1,87	Hs.750	fibrillin 1 (Marfan
							syndrome)
1231	GAAGCAATAAA	1,67	6	3,59	1,87	Hs.198253	major histocompatibility
							complex, class II, DQ
	070111070	12.2					alpha
1232	GTGAAACTCCG	10,34	37	3,58	9,36	Hs.261734	ESTs, Moderately similar
4000	00070000407	0.50		0.55			to ALU7_HUMAN ALU
1233	GGCTGCCGAGT	0,56	2	3,57	0,72	Hs.99829	hypothetical protein
1004	CTTACCCCCAC	0.50		0.57	0.70	11- 00000	FLJ20565
	CTTAGCCCCAG	0,56	2	3,57		Hs.96908	ESTs
1235	TTATTCCACAA	0,56	2	3,57	0,72	Hs.93765	lipoma HMGIC fusion
1226	TCACAGCCCCC	0,56		3,57	0.70	Hs.8619	partner
1230	I ONONGCOCCO	0,50		3,5/	0,72	118.0019	SRY (sex determining
1237	TTTTCCTTTTG	0,56	2	3,57	0.72	Hs.78546	region Y)-box 18 ATPase, Ca++
1237		0,56	2	3,51	0,72	⊓S.70040	
							transporting, plasma membrane 1
1238	CTTGCATAAGA	0,56		3,57	0.72	Hs.72912	cytochrome P450,
'200	STICOMINACA .	0,55		ا ''د,ب	0,12	13.7 23 12	subfamily I (aromatic
							compound-indu
1239	TATGTGTTCTC	0,56	2	3,57	0.72	Hs.3353	beta-1,3-
		-,00	_	ا ''' ا	٥, ، ح		glucuronyltransferase 1
		ļ i					(glucuronosyltransf
1240	ACTATCATCTT	0,56	2	3,57	0.72	Hs.29117	H.sapiens mRNA for pur
1		'	_	-,	-,		alpha extended
1							3'untranslated
						-	•

1241	CGCCTATAGTC	0,56	2	3,57	0.72	Hs.271166	ESTs, Moderately similar
							to ALU7_HUMAN ALU
	CCCGCCAGTGC	0,56		3,57	0,72	Hs.256297	integrin, alpha 11
1243	TTCTAATTTTT	0,56	2	3,57	. 0,72	Hs.170414	paired basic amino acid
		ļ	_				cleaving system 4
	CGGGAAGACAT	0,56		3,57	0,72	Hs.154525	KIAA1076 protein
	CCAGTAGTCCC	0,56	2	3,57	0,72	Hs.147959	EST
<u> </u>	CAGTTTGTGTT	0,56	2	3,57	0,72	Hs.144477	hypothetical protein PRO2975
	TCCTTTAAAAT	0,56		3,57		Hs.10587	KIAA0353 protein
	GTGCTAAGCGG	12,92		3,56		Hs.4217	collagen, type VI, alpha 2
	AGAATCACTTG	11,6		3,53		Hs.117582	CGI-43 protein
	GTGGTGTACGC	2,55	9	3,53		Hs.182225	RNA binding motif protein 3
1251	GCCCCAGAATC	0,85	3	3,53	1,02	Hs.8682	ESTs, Moderately similar to ALU1_HUMAN ALU
1252	TTGGGAGGCTG	0,85	3	3,53	1,02	Hs.118269	ESTs, Weakly similar to A46010 X-linked retinopathy
1253	GTGGCACGCGC	5,39	19	3,53	5,01	Hs.187346	ESTs
	CAAGCGCTCTA	1,14		3,51		Hs.23598	CREB binding protein
1							(Rubinstein-Taybi syndrome)
1255	AGTTCGAGACC	1,14	4	3,51	1,29	Hs.232540	ESTs
1256	AGAACCAAAAA	1,14		3,51	1,29	Hs.181244	major histocompatibility
	, ,						complex, class I, A
1257	CTGGCTATCCG	1,14	4	3,51	1,29	Hs.10784	hypothetical protein FLJ20037
1258	TAGTCCCAGCT	3,14	11	3,50	3,08	Hs.274579	ancient conserved domain protein 1
1259	GTGAAATCCTG	8	28	3,50	7,06	Hs.53531	lipoic acid synthetase
1260	CCTGTAATTCC	13,48	47	3,49	11,36	Hs.23582	tumor-associated calcium signal transducer 2
	CTTCTTGCCCC	5,74	20	3,48	5,17	Hs.251577	hemoglobin, alpha 1
	TGGTTGGTGGT	3,16		3,48		Hs.12701	plasmolipin
	CCCGTAATCCC	4,89	17	3,48	4,46	Hs.274168	Homo sapiens mRNA; cDNA DKFZp761P0212 (from
1264	CCTGGCCAGAA	1,44	5	3,47	1,55	Hs.261734	ESTs, Moderately similar to ALU7_HUMAN ALU
	CGTGTAATCCC	3,46		3,47		Hs.187761	ESTs
	GCGAAACCTCA	1,73	6	3,47	1,81	Hs.42644	thioredoxin-like
1267	TGGTTACAAAA	1,73	6	3,47	1,81	Hs.3850	Homo sapiens clone 23596 mRNA sequence
	CCACAGCACTC	2,02	7	3,47	2,06	Hs.273828	ESTs
1269	GTGGCACGTGC	29,86	103	3,45	23,64	Hs.278588	ESTs, Weakly similar to ALU2_HUMAN ALU
1270	ATGTCTTTCT	5,22	18	3,45	4,66	Hs.1516	insulin-like growth factor- binding protein 4
1271	CTTGTAGTCCC	3,19	11	3,45	3,02	Hs.272202	hypothetical protein

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10==	07047070	-		-			FLJ20825
	CTCATCTGCTG	2,9		3,45		Hs.82109	syndecan 1
	AGGACATAATT	0,29		3,45		Hs.93454	ESTs
1274	GAGCTACACCA	0,29	1	3,45	0,34	Hs.82171	Human clone 191B7
ĺ		i					placenta expressed mRNA
L							from
	TTGCTACTAAA	0,29	1			Hs.7790	ESTs
1276	CTTAGTGTTTT	0,29	1	3,45	0,34	Hs.7720	dynein, cytoplasmic, heavy
							polypeptide 1
	CTGGTCCTGGA	0,29		3,45	0,34	Hs.76476	cathepsin H
1278	GGTGGCAGTTG	0,29	1	3,45	0,34	Hs.75794	endothelial differentiation,
							lysophosphatidic acid G
1279	CTGATATAGAC	0,29	1	3,45	0,34	Hs.74002	nuclear receptor
							coactivator 1
	TGGAAATCATT	0,29		3,45		Hs.5028	DKFZP564O0423 protein
1281	ACTTTGAAAGG	0,29	1	3,45	0,34	Hs.44077	hypothetical protein
							FLJ10793
	AAAGGCACTGA	0,29		3,45		Hs.3994	ESTs
1283	GTTCTCTTTTT	0,29	1	3,45	0,34	Hs.3843	dual specificity
<u>.</u> '							phosphatase 7
1284	AGCGCAGCTGT	0,29		3,45		Hs.34771	ESTs
	ATTGTGAAGAG	0,29		3,45		Hs.34578	alpha2,3-sialyltransferase
1286	GCCTTCGGAAA	0,29	1	3,45	0,34	Hs.33104	Homo sapiens mRNA;
							cDNA DKFZp434H2121
							(from
	GCTATTTTGAT	0,29		3,45		Hs.32250	ESTs
	TATCTCTTAAA	0,29		3,45	0,34	Hs.286228	ESTs
1289	TGTGATTTTTA	0,29		3,45		Hs.286163	ESTs
	TATTTCAGATT	0,29	1	3,45	0,34	Hs.285585	ESTs
1291	ATGATTTTGAG	0,29	1	3,45	0,34	Hs.285306	putative selenocysteine
							lyase
1292	AATATTCATAT	0,29	1	3,45	0,34	Hs.284311	Homo sapiens clone
l						<u>.</u>	25038 mRNA sequence
	AGACCCCATTT	0,29	_1	3,45		Hs.279297	EST
1294	CACCCATAGTC	0,29	1	3,45	0,34	Hs.278018	EST
	GTCTTGCTGCA	0,29	1	3,45	0,34	Hs.26966	KIAA1171 protein
1296	CGGCCCATCTG	0,29	1	3,45	0,34	Hs.26290	ESTs
	CAATCTGATGC	0,29	1	3,45		Hs.26176	hypothetical protein
							FLJ10261
1298	ATGTTGGGTGT	0,29	1	3,45	0,34	Hs.260855	Homo sapiens mRNA;
					·		cDNA DKFZp761G2311
							(from
1299	ACTCTGTCTCC	0,29	1	3,45	0,34	Hs.259339	EST
1300	GGAATACAGAA	0,29	1	3,45		Hs.250825	ESTs, Highly similar to
					•		vacuolar protein sorting
							hom
1301	ACTGGGCAAGC	0,29		3,45		Hs.240062	hypothetical protein
1302	GGTCAGAAATT	0,29	1	3,45	0,34	Hs.211581	metal-regulatory
							transcription factor 1
1303	AACCGAAGGGA	0,29	1	3,45	0,34	Hs.20596	ESTs

1304	GGGATAGAGAC	0,29	1	3,45	0,34	Hs.202955	hypothetical protein FLJ20507
1305	TTTCAGTTAGT	0,29	1	3,45	0,34	Hs.196284	ESTs
1306	GCTATTGATGT	0,29	1	3,45	0,34	Hs.193398	ESTs
1307	AAGAGGAGGCC	0,29	1	3,45	0,34	Hs.183639	hypothetical protein FLJ10210
1308	TAATACACTAA	0,29	1	3,45	0,34	Hs.183475	Homo sapiens clone 25061 mRNA sequence
1309	GCAGATGCTTT	0,29	1	3,45	0,34	Hs.180799	ESTs, Moderately similar to ALU5_HUMAN ALU
1310	TCACAAGGCTG	0,29	1	3,45	0,34	Hs.17998	ESTs
1311	TGCGAGCTGGG	0,29	1	3,45	0,34	Hs.179573	collagen, type I, alpha 2
1312	GGATTTGCTGC	0,29	1	3,45	0,34	Hs.177956	Homo sapiens mRNA; cDNA DKFZp434C0926 (from
1313	CACGCACACAC	0,29	1	3,45	0,34	Hs.177664	KIAA0914 gene product
1314	ATGTGGACTGA	0,29	1	3,45		Hs.174905	KIAA0033 protein
1315	GGAGGCAGAGC	0,29	1	3,45	0,34	Hs.172838	Human clone Z'3-1 placenta expressed mRNA from
1316	GCTTTCTGTAA	0,29	1	3,45	0,34	Hs.172674	nuclear factor of activated T-cells, cytoplasmic 3
	TGATTATTTAC	0,29	1	3,45	0,34	Hs.16930	ESTs
1318	ACATCTGCCTG	0,29	1	3,45	0,34	Hs.161554	hypothetical protein FLJ20159
1319	CTTAGTTTTAA	0,29	1	3,45	0,34	Hs.161554	hypothetical protein FLJ20159
1320	AGGAAGAGTCA	0,29	1	3,45	0,34	Hs.154655	imogen 38
1321	CAATGCAGAGG	0,29	1	3,45	0,34	Hs.150748	malonyl-CoA decarboxylase
	TAATTCTTGTA	0,29	1	3,45	0,34	Hs.146123	ESTs
1323	CAAGGGCCCAC	0,29	1	3,45	0,34	Hs.14587	ESTs, Weakly similar to AF151859_1 CGI-101 protein [
1324	TTTTGAAGAAA	0,29	1	3,45	0.34	Hs.144465	ESTs
	TGTCTCCGTCT	0,29	1	3,45		Hs.135150	lung type-I cell membrane- associated glycoprotein
	TTTTCTTCAGG	0,29	1	3,45	0,34	Hs.125753	chromosome 2 open reading frame 2
1327	TTCCTCCCTCT	0,29	1	3,45	0,34	Hs.125384	ESTs
	TCTGCCTTTCT	0,29		3,45		Hs.125019	ESTs, Highly similar to KIAA0886 protein [H.sapiens]
	ATAACTGTCAG	0,29		3,45	0,34	Hs.12040	STE20-like kinase
1330	TTGCAGTTTTT	0,29		3,45		Hs.117582	CGI-43 protein
	TATTTAAAAAA	0,29	1	3,45	0,34	Hs.117304	ESTs
L	GGCTCAGGGGC	0,29	1	3,45		Hs.116489	ESTs, Weakly similar to GCP170 [H.sapiens]
	TTTATTGAAAC	0,29	1	3,45	0,34	Hs.112193	mutS (E. coli) homolog 5
1334	CTGGCTTAAAT	0,29	1	3,45		Hs.11171	APG5 (autophagy 5, S.
		-					

		<del> </del>				<del>,</del>	
122							cerevisiae)-like
1335	GCTTTATGTGG	0,29	1	3,45	0,34	Hs.111460	ESTs, Weakly similar to Con1 [H.sapiens]
1336	GCATACTTTAT	0,29	1	3,45	0.34	Hs.109370	ESTs
	AGAATACTGAG	0,29			0.34	Hs.106705	neuronal PAS domain
							protein 2
1338	ACCCAAAAAA	0,29	1	3,45	0,34	Hs.101840	major histocompatibility
							complex, class I-like seque
1339	GGACATTAGGG	0,29	1	3,45	0,34	Hs.101265	MEMOREC NSM2 (CCA1) sphingomyeli
1340	AAAAATAAATT	0,29	1	3,45	0,34	Hs.100261	Homo sapiens mRNA; cDNA DKFZp564B222 (from clone
1341	AGATACATAGC	1,46	5	3,42	1.53	Hs.84045	Homo sapiens cDNA
		.,			,,00		FLJ20288 fis, clone HEP04414
1342	TGGATATCAGT	1,46	5	3,42	1,53	Hs.7327	claudin 1
	TTTTCCACTTT	1,46		3,42		Hs.6900	ring finger protein 13
	GTGGCTCAGGC	1,17	4	3,42	1,26	Hs.259047	<b>ESTs</b>
	CCTGTGATTCC	1,17	4	3,42	1.26	Hs.227961	EST
	GGCGACAGAGC	3,22		3,42		Hs.92254	hypothetical protein FLJ20163
1347	сствтветсст	3,22	11	3,42	2,99	Hs.120769	Homo sapiens cDNA FLJ20463 fis, clone KAT06143
1348	TGCCTGTGGTC	4,69	16	3,41	4,15	Hs.277100	ESTs, Weakly similar to ALU2_HUMAN ALU
1349	GTAAAAAAGCC	0,88	3	3,41	0,99	Hs.98988	ESTs
1350	TGTGAACACAT	0,88	3	3,41	0,99	Hs.80645	interferon regulatory factor
1351	AAACGAAGTTG	0,88	3	3,41	0,99	Hs.78353	SFRS protein kinase 2
1352	TACATCAGTAA	0,88	3	3,41	0,99	Hs.65029	growth arrest-specific 1
1353	CCTGTAGGCCC	0,88	3	3,41	0,99	Hs.207938	EST
1354	GTGAGACCTCG	0,88	3	3,41	0,99	Hs.203206	ESTs, Moderately similar to ALU1_HUMAN ALU
1355	TGCCACCACAC	2,64	9	3,41	2,51	Hs.239993	ESTs
	GCTGGATTTTG	0,59	2	3,39		Hs.82124	laminin, beta 1
1357	TCACTTTTTA	0,59	2	3,39	0,69	Hs.8045	ESTs
	ATTATCCTCAG	0,59	2	3,39	0.69	Hs.7987	DKFZP434F162 protein
	GGATCCAATTT	0,59		3,39		Hs.61796	transcription factor AP-2
		-,00		5,00	2,00		gamma (activating lenhancer
1360	CCAATTGAAGA	0,59	2	3,39	0,69	Hs.40328	ESTs
	TTACTTTTGGT	0,59	2	3,39	0,69	Hs.285861	hypothetical protein FLJ10359
1362	GAGAGCTTTGC	0,59		3,39	,	Hs.275374	aldo-keto reductase family 1, member C1 (dihydrodiol
1363	TACCCCCAAAC	0,59	2	3,39	0,69	Hs.241926	ESTs
1364	GGGCAGACACT	0,59	2	3,39	0,69	Hs.18878	ESTs, Weakly similar to dJ876B10.4 [H.sapiens]

1365	ATGGCGCACGC	0,59	2	3,39	0,69	Hs.124984	ESTs, Moderately similar
							to unnamed protein
		l					product
1366	AGGTTGCCGAG .	0,59	2	3,39	0,69	Hs.105399	KIAA0809 protein
1367	CCACTGCACCC	5,92		3,38		Hs.6853	carbohydrate (N-
1		1 1		'			acetylglucosamine 6-O)
							sulfotransfer
1368	ATGGTGGGGA	5,33	18	3,38	4,55	Hs.1665	zinc finger protein
	[				·		homologous to Zfp-36 in
							mouse
1369	CCCTCTCCCTT	2,37	8	3,38	2,24	Hs.85087	latent transforming growth
		'					factor beta binding prote
1370	TCACCAAAAA	1,79	6	3,35	1,75	Hs.84753	KIAA0246 protein
	GTGAAACCCCC	17,64				Hs.265865	EST
	CCTGCAATCCC	10,2		3,33		Hs.3280	caspase 6, apoptosis-
				-,-,-	.,		related cysteine protease
1373	GTGAAGCCCCG	6,62	22	3,32	5.34	Hs.285592	Homo sapiens mRNA;
		5,5-		,,,,	-,		cDNA DKFZp564M113
		l					(from
1374	CCACTGTACTC	16	53	3,31	11.94	Hs.220261	ESTs, Moderately similar
				0,0.	,.		to ALU4 HUMAN ALU
1375	GTGGTGGGCGC	15,12	50	3,31	11.28	Hs.136810	ESTs, Weakly similar to
		,					ALU1 HUMAN ALU
1376	AGTATGACCTA	0,91	3	3,30	0.96	Hs.74649	cytochrome c oxidase
1		-,-		-,			subunit VIc
1377	GTGACAGCCAC	0,91	3	3,30	0,96	Hs.74441	chromodomain helicase
					·		DNA binding protein 4
1378	GGGCTTTTGAG	0,91	3	3,30	0,96	Hs.29893	Homo sapiens mRNA full
				Ċ			length insert cDNA clone
						•	EURO
1379	GTGAGACCCCT	0,91	3	3,30	0,96	Hs.269952	ESTs, Weakly similar to
							ALU1_HUMAN ALU
1380	GTGGTGCACAT	0,91	3	3,30	0,96	Hs.269030	ESTs
1381	CCTGTAGTCAC	0,91	3	3,30	0,96	Hs.268900	ESTs
1382	TGGTAACTGGC	0,91	3	3,30	0,96	Hs.108741	ESTs
1383	GTGGTATGTGC	1,52		3,29		Hs.277102	ESTs, Weakly similar to
					·		ALU1_HUMAN ALU
1384	GTAAGATTAGC	1,52	5	3,29	1,47	Hs.250705	ESTs
	GCGAAACCCCA	21,97		3,28		Hs.210682	ESTs, Weakly similar to
							ALU6_HUMAN ALU
1386	ATCGTGCCACT	6,12	20	3,27	4,81	Hs.7615	Homo sapiens mRNA;
		]			•		cDNA DKFZp434N2030
L							(from
1387	TCTGTAATCCC	13,48	44	3,26	9,85	Hs.142	sulfotransferase family,
					•		cytosolic, 1A, phenol-prefe
1388	TTAGCCAGGCT	3,37	11	3,26	2,85	Hs.71367	ESTs, Moderately similar
		] _ ` }					to ALU7_HUMAN ALU
1389	ACAAAACCCTG	1,23	4	3,25	1,21	Hs.268591	ESTs
	ATCTCGGCTCA	4,31		3,25		Hs.29809	Homo sapiens mRNA;
		[					cDNA DKFZp434C185

1391 CCTGTAATGCC         4,01         13 3,24         3,26 Hs.7179         RAD homo homo homo homo homo homo homo hom	, Moderately similar U4_HUMAN ALU D sapiens mRNA; A DKFZp434G2127  , Weakly similar to C HUMAN !!!! ALU SS Weakly similar to G HUMAN ALU FAMILY kin A4 xylesterase 1
1392   CCACCGCACTC   6,8   22   3,24   5,18   Hs.222669   ESTS to AL	Noderately similar U4_HUMAN ALU D sapiens mRNA; A DKFZp434G2127  Weakly similar to HUMAN IIII ALU SS Weakly similar to HUMAN ALU FAMILY kin A4 xylesterase 1
1392         CCACCGCACTC         6,8         22         3,24         5,18         Hs.222669         ESTs to AL t	, Moderately similar U4_HUMAN ALU D sapiens mRNA; A DKFZp434G2127  , Weakly similar to C HUMAN !!!! ALU SS Weakly similar to G HUMAN ALU FAMILY kin A4 xylesterase 1
to AL 1393 GTGGTGTGC 11,75 38 3,23 8,5 Hs.27038 Homo cDNA (from 1394 ATGAAACCCCA 8,35 27 3,23 6,22 Hs.285341 ESTs ALUC CLAS 1395 CCTGTAGCCCC 2,17 7 3,23 1,92 Hs.277320 EST, ALU6 SUBF 1396 TTTTTAAAAAA 0,62 2 3,23 0,66 Hs.77840 annex 1397 AAGGAGCAAGT 0,62 2 3,23 0,66 Hs.76688 carbo (mone) serine	U4_HUMAN ALU D sapiens mRNA; A DKFZp434G2127  , Weakly similar to C HUMAN !!!! ALU DS Weakly similar to C HUMAN ALU FAMILY kin A4 xylesterase 1
1393         GTGGTGTGC         11,75         38         3,23         8,5         Hs.27038         Homo cDNA (from (from (from 1394))           1394         ATGAAACCCCA         8,35         27         3,23         6,22         Hs.285341         ESTs ALUC CLAS           1395         CCTGTAGCCCC         2,17         7         3,23         1,92         Hs.277320         EST, ALU6 SUBF           1396         TTTTTAAAAAA         0,62         2         3,23         0,66         Hs.77840         annex           1397         AAGGAGCAAGT         0,62         2         3,23         0,66         Hs.76688         carbo (mond sering)	o sapiens mRNA; A DKFZp434G2127  , Weakly similar to C HUMAN !!!! ALU SS  Weakly similar to C HUMAN ALU FAMILY kin A4  xylesterase 1
CDNA	N DKFZp434G2127  , Weakly similar to CHUMAN IIII ALUSS  Weakly similar to CHUMAN ALUSE HUMAN ALUSE FAMILY AND ALUSE
1394 ATGAAACCCCA	, Weakly similar to LHUMAN IIII ALU S Weakly similar to LHUMAN ALU FAMILY kin A4 xylesterase 1
1394         ATGAAACCCCA         8,35         27         3,23         6,22         Hs.285341         ESTs ALUC CLAS           1395         CCTGTAGCCCC         2,17         7         3,23         1,92         Hs.277320         EST, ALU6 SUBF           1396         TTTTTAAAAAA         0,62         2         3,23         0,66         Hs.77840         annex           1397         AAGGAGCAAGT         0,62         2         3,23         0,66         Hs.76688         carbo (money sering)	, Weakly similar to C_HUMAN !!!! ALU SS Weakly similar to C_HUMAN ALU FAMILY kin A4 xylesterase 1
ALUC CLAS  1395 CCTGTAGCCCC 2,17 7 3,23 1,92 Hs.277320 EST, ALU6 SUBF  1396 TTTTTAAAAAA 0,62 2 3,23 0,66 Hs.77840 annex 1397 AAGGAGCAAGT 0,62 2 3,23 0,66 Hs.76688 carbo (mone serine	HUMAN IIII ALU SS Weakly similar to HUMAN ALU FAMILY kin A4 xylesterase 1
CLAS   1395   CCTGTAGCCCC   2,17   7   3,23   1,92   Hs.277320   EST, ALU6   SUBF   1396   TTTTTAAAAAA   0,62   2   3,23   0,66   Hs.77840   annex   1397   AAGGAGCAAGT   0,62   2   3,23   0,66   Hs.76688   carbo (mone serine	SS Weakly similar to LHUMAN ALU AMILY Kin A4 xylesterase 1
1395         CCTGTAGCCCC         2,17         7         3,23         1,92         Hs.277320         EST, ALU6 SUBF           1396         TTTTTAAAAAA         0,62         2         3,23         0,66         Hs.77840         annex           1397         AAGGAGCAAGT         0,62         2         3,23         0,66         Hs.76688         carbo (mond serine)	Weakly similar to LHUMAN ALU FAMILY kin A4 xylesterase 1
ALU6 SUBF  1396 TTTTTAAAAAA 0,62 2 3,23 0,66 Hs.77840 annex 1397 AAGGAGCAAGT 0,62 2 3,23 0,66 Hs.76688 carbo (mone serine	LHUMAN ALU FAMILY kin A4 xylesterase 1
SUBF   1396   TTTTTAAAAAA   0,62   2 3,23   0,66   Hs.77840   annex   1397   AAGGAGCAAGT   0,62   2 3,23   0,66   Hs.76688   carbo (mone serine	-AMILY kin A4 xylesterase 1
1396 TTTTTAAAAAA 0,62 2 3,23 0,66 Hs.77840 annex 1397 AAGGAGCAAGT 0,62 2 3,23 0,66 Hs.76688 carbo (monts) serine	kin A4 xylesterase 1
1397 AAGGAGCAAGT 0,62 2 3,23 0,66 Hs.76688 carbo (mono serine	xylesterase 1
(mone serine	
<u></u>	ocyte/macrophage
1398 ACTITITATG 0.62 2 3 23 0 66 Hs 697 cytoc	e ester
	hrome c-1
1399 ATTGAGCCACA 0,62 2 3,23 0,66 Hs.63290 2-hyd	roxyphytanoyl-CoA
lyase	
	nate dehydrogenase
	lex, subunit A,
flavor	
1401 ATCACAGCTCA 0,62 2 3,23 0,66 Hs.29590 ESTs	
	, Weakly similar to
	atively spliced
produ	
	sapiens cDNA
	1265 fis, clone
	E1009158
	modulin 3
1406 ATGATAATTAA 0,62 2 3,23 0,66 Hs.170142 ESTs	uitous)
	er (mouse) maf-
	d leucine zipper
homo	
1408 GAGAGAGAA 0,62 2 3,23 0,66 Hs.169391 ESTs	
1409 TACCCTGAAAC 0,62 2 3,23 0,66 Hs.144018 ESTs	
	cellular matrix protein
1 1 2 2 2 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3	Johann Hank protein
1411 GGTGAGCGTGT 1,55 5 3,23 1,44 Hs.2913 EphB	3
1412 GTGGTGGATGC 1,55 5 3,23 1,44 Hs.277904 EST	
	hetical protein
	, Weakly similar to
	HUMAN IIII ALU
1415 CCCACTTGTAA 3,75 12 3,20 3 Hs.75922 brain	protein I3

1416	GCCCTTTCTCT	4,07	13	3,19	3,21	Hs.7835	endocytic receptor
							(macrophage mannose receptor fami
1417	AGACCTCCTTC	1,88	6	3,19	1,67	Hs.281706	sortilin 1
1418	AGTGGTGGCTA	1,88		3,19		Hs.230	fibromodulin
	GGACAGATGTA	0,94	3	3,19	0,93	Hs.75356	transcription factor 4
1420	GTGGCGAGCAC	0,94	3	3,19	0,93	Hs.261831	EST
	ATGGTGTGTGC	0,94	3	3,19	0,93	Hs.193347	ESTs
	GGACTGAGTCA	0,94	3	3,19	0,93	Hs.18387	transcription factor AP-2 alpha (activating enhancer
1423	GTGAGTGCCCT	0,94	3	3,19	0,93	Hs.171872	DEAD/H (Asp-Glu-Ala- Asp/His) box polypeptide 8 (RNA
1424	GCGGAACCTCA	0,94	3	3,19	0,93	Hs.10700	hypothetical protein
1425	GTGTGGGGGGC	12,86		3,19		Hs.2340	junction plakoglobin
	GTGAAACTCCA	10,05		3,18		Hs.140002	ESTs, Moderately similar to ALU7_HUMAN ALU
	CCACTACACTC	9,11		3,18	·	Hs.83429	tumor necrosis factor (ligand) superfamily, member 1
	ACGGAAGTTTT	1,26		3,17	1,18	Hs.144974	ESTs, Highly similar to unnamed protein product [H.s
	GTGAAACCCGT	4,1	13	3,17	3,18	Hs.278577	Homo sapiens mRNA; cDNA DKFZp564P073 (from clone
1430	TCAAACTGTGA	1,58		3,16	1,42	Hs.94881	ESTs
	CATCGAAAGTT	0,64	2	3,13	0,64	Hs.80618	hypothetical protein
1432	AGTAATCATCA	0,64	2	3,13	0,64	Hs.75925	proteasome (prosome, macropain) inhibitor subunit 1
	AATAATCCTGG	0,64	2	3,13	0,64	Hs.62908	ESTs
	GTATTCCTAAA	0,64	2	3,13	0,64	Hs.5724	ESTs, Weakly similar to multi PDZ domain protein MUP
	CTGGGAAGCAT	0,64	2	3,13	0,64	Hs.42311	ESTs
	GATCAAAACTG	0,64	2	3,13	0,64	Hs.41267	c21orf7 form A-D
	GTGACAGGCGC	0,64	· 2		0,64	Hs.278879	ESTs, Moderately similar to ALUA_HUMAN !!!! ALU
· .	GCACCGTGGAA	0,64		3,13	0,64	Hs.27299	transcriptional regulator protein
	TTAACTGTATT	0,64		3,13		Hs.180952	actin, beta
	ATGTTAGAGAC	0,64	2	3,13	0,64	Hs.1592	CDC16 (cell division cycle 16, S. cerevisiae, homolo
	ATCGCATCACT	0,64		3,13		Hs.158126	ESTs
	GACTCTGGAGA	0,64		3,13	_	Hs.154567	supervillin
	AAACTGTTCAA	0,64		3,13		Hs.118978	KIAA0256 gene product
	ACCAACACGGG	0,64		3,13		Hs.109005	ESTs
1445	AAAGATGTATC	0,32	1	3,13	0,32	Hs.979	pyruvate dehydrogenase (lipoamide) beta

1446	AAAACAGCAAG	0,32	1	3 13	0.32	Hs.92909	SON DNA binding protein
	TTTTCAGGTAA	0,32	1	3,13 3,13	0,32	Hs.91773	protein phosphatase 2
1777	ITTTOAGGTAA	0,52	•	3, 13	0,52	113.91770	(formerly 2A), catalytic
							subun
1448	TGCCTCCCAGC	0,32	1	3,13	0.32	Hs.90527	HSPC128 protein
	TAAGTGAACAT	0,32		3,13		Hs.83164	collagen, type XV, alpha 1
	TTTTGCTCAGA	0,32		3,13		Hs.8102	ribosomal protein S20
	CATTCTCCCAG	0,32	1	3,13		Hs.79110	nucleolin
1452	GTTTCAGCACT	0,32	1	3,13	0,32	Hs.77502	methionine
							adenosyltransferase II,
					. !		alpha
1453	GTAACTCTATG	0,32	1	3,13	0,32	Hs.7277	peroxisomal biogenesis
							factor 3
	GTTCTATTGTA	0,32		3,13		Hs.6909	DKFZP564G202 protein
	CTATATTGTAA	0,32	1	3,13		Hs.65919	ESTs
1456	GTGAAACATTG	0,32	1	3,13	0,32	Hs.6567	Homo sapiens mRNA;
						34	cDNA DKFZp434C136
1457	GTATTGAAGTT	0,32	1	3,13	0.22	Hs.6079	(from clone  B cell RAG associated
1457	GIAIIGAAGII	0,32	1:	3,13	0,32	ns.60/9	protein
1/58	CTTTAGAAGCA	0,32	1	3,13	0.32	Hs.5669	ESTs
	TGACTCCTCAA	0,32	1	3,13		Hs.47007	mitogen-activated protein
1400	10/010010/4	0,02	•	0,10	0,02	113.47 007	kinase kinase kinase 14
1460	CTTTTATGGAC	0,32	1	3,13	0.32	Hs.44833	ESTs
	TAAATCTACAA	0,32	1.	3,13		Hs.44701	ESTs
	TTCCCAAATGA	0,32	1	3,13	0.32	Hs.44257	Homo sapiens mRNA;
	,						cDNA DKFZp762O2215
							(from
	GTAAGAGTTCT	0,32	1	3,13	0,32	Hs.4084	KIAA1025 protein
1464	ATGCCATTGGA	0,32	1	3,13	0,32	Hs.30213	ceroid-lipofuscinosis,
							neuronal 5
	AGGAAATGGAT	0,32	·1	3,13 3,13	0,32	Hs.30194	ESTs
1466	AACAAGCTGGG	0,32	1	3,13	0,32	Hs.29759	RNA POLYMERASE I
							AND TRANSCRIPT
4407	TOOAOTTOAOA	0.00		0.40	0.00	11- 00055	RELEASE
	TGCACTTGAGA GCATTCTGGTT	0,32	<u>1</u>	3,13 3,13	0,32	Hs.29055	ESTs
						Hs.286261	ESTs
	GTGGCCACCCT AAGGTGGTTGT	0,32		3,13		Hs.286217	KIAA0685 gene product
14/0	~~001001101	0,32	. 1	3,13	0,32	Hs.285999	trinucleotide repeat containing 15
1471	AGAACTACGTG	0,32	1	3,13	0.33	Hs.284176	hypothetical protein
177	, .5, , , , , , , , , , , , , , , , , ,	0,02	•	اک، اکا	0,02	5.207170	PRO2221
1472	AATTTGGGAGA	0,32	1	3,13	0.32	Hs.279882	PC326 protein
	GGGAAACCCCT	0,32		3,13		Hs.279408	EST
	CTTACTCTTGA	0,32		3,13		Hs.27342	ESTs
	ACCGTGCCACT	0,32		3,13		Hs.270667	ESTs
	AGGCTGGTTTA	0,32	1			Hs.26322	cell cycle related kinase
	TCTTTTGGGAG	0,32		3,13		Hs.257312	EST
1478	ACTGATCTTGT	0,32		3,13		Hs.251871	CTP synthase

1480   ATGGGGAAAGA   0,32					-			1-00
1481   ACAGTGCCACT   0,32			0,32					ESTs
1482   TGTGGGGACAA   0,32							Hs.24989	
1483   CAAGTCTCCAG	1481	ACAGTGCCACT	0,32	1	3,13	0,32	Hs.246374	ESTs
1483   CAAGTCTCCAG	1482	TGTGGGGACAA	0,32	1	3,13	0,32	Hs.245017	EST
1484   TTCCCTCCAAA   0,32	1483	CAAGTCTCCAG	0.32	1	3.13	0.32	Hs.241515	COX11 (yeast) homolog.
assembly   assembly					,	-,		
1484   TTCCCTCAAA   0,32			J			ļ	, · .	
1485   ATGTATAGGGC   0,32	1484	TTCCCTCCAAA	0.32	1	3 13	0.32	Hs 239727	
1486   GGTTGTTGCGG								
1487   TCTTCTTAATA								
1488   GCTCATTTCAG   0,32								
1488   GCTCATTTCAG	1407	ICHCHAAIA	0,32	, 	3, 13	0,32		predicted using
Length insert cDNA clone   Length   L	4400	COTOATTTOAC	0.00		0.40	0.00		
1489   AGTTCCAGACC   0,32	1488	GCTCATTTCAG	0,32	1	3,13	0,32	Hs.22870	
1489 AGTTCCAGACC         0,32         1 3,13         0,32 Hs.223935         EST           1490 AGTCAGTGGGA         0,32         1 3,13         0,32 Hs.21943         ESTs, Weakly similar to ORF YGL221c [S.cerevisiae]           1491 TTTCCAATGGA         0,32         1 3,13         0,32 Hs.21756         translation factor suif homolog           1492 TACATTTGAAT         0,32         1 3,13         0,32 Hs.21537         protein phosphatase 1, catalytic subunit, beta isofo           1493 TTTCTGTATT         0,32         1 3,13         0,32 Hs.21557         hypothetical protein DKFZp762K2015           1494 AGAAAGATGGA         0,32         1 3,13         0,32 Hs.211577         kinectin 1 (kinesin receptor)           1495 TTTACCTTTGG         0,32         1 3,13         0,32 Hs.21108         ESTs           1496 TACGATATTCA         0,32         1 3,13         0,32 Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1497 GCACTGGGGCA         0,32         1 3,13         0,32 Hs.197751         KIAA0666 protein           1498 GCTGCTAGAAA         0,32         1 3,13         0,32 Hs.196437         hypothetical protein FLJ1078           1500 ATCGGCTCCCA         0,32         1 3,13         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32	- · · ·							
1490   AGTCAGTGGGA   0,32								
1491   TTTCCAATGGA								
1491 TTTCCAATGGA	1490	AGTCAGTGGGA	0,32	1	3,13	0,32	Hs.21943	
1491         TTTCCAATGGA         0,32         1         3,13         0,32 Hs.21756         translation factor sui1 homolog           1492         TACATTTGAAT         0,32         1         3,13         0,32 Hs.21537         protein phosphatase 1, catalytic subunit, beta isofo           1493         TTTTCTGTATT         0,32         1         3,13         0,32 Hs.21356         hypothetical protein DKFZp762K2015           1494         AGAAAGATGGA         0,32         1         3,13         0,32 Hs.211577         kinectin 1 (kinesin receptor)           1495         TTTACCTTTGG         0,32         1         3,13         0,32 Hs.2108         ESTs           1496         TACGATATTCA         0,32         1         3,13         0,32 Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1498         GCTGCTAGAAA         0,32         1         3,13         0,32 Hs.197751         KIAA0666 protein pDKFZpd34O1572 (from FLJ10788           1500         ATCGGCTCCCA         0,32         1         3,13         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from FLJ10788           1501         ATAATGGAGTG         0,32         1         3,13         0,32 Hs.172506         myosin VB           1503         TGCCTGACAAG         0,32								
1492   TACATTTGAAT   0,32	1491	TTTCCAATGGA	0.32	1	3 13	0.32	He 21756	
1492         TACATTTGAAT         0,32         1         3,13         0,32         Hs.21537         protein phosphatase 1, catalytic subunit, beta isofo           1493         TTTTCTGTATT         0,32         1         3,13         0,32         Hs.21356         hypothetical protein DKFZp762K2015           1494         AGAAAGATGGA         0,32         1         3,13         0,32         Hs.211577         kinectin 1 (kinesin receptor)           1495         TTTACCTTTGG         0,32         1         3,13         0,32         Hs.21108         ESTs           1496         TACGATATTCA         0,32         1         3,13         0,32         Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1498         GCTGCTAGAAA         0,32         1         3,13         0,32         Hs.197751         KIAA0666 protein           1499         GGGTAGAGAGAT         0,32         1         3,13         0,32         Hs.196437         hypothetical protein FLJ10788           1500         ATCGGCTCCCA         0,32         1         3,13         0,32         Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501         ATAATGGAGTG         0,32         1         3,13         0,32         Hs.172506         myos		11100/1100/1	0,02	•	0, 10	0,02	113.21750	
Catalytic subunit, beta isofo	1492	TACATTTGAAT	0.33	1	2 12	0.33	Uc 21527	
1493   TTTTCTGTATT   0,32	1432	INCATTION	0,32	'	3,13	0,32	П8.2 155 <i>1</i>	
1493         TTTTCTGTATT         0,32         1         3,13         0,32         Hs.21356         hypothetical protein DKFZp762K2015           1494         AGAAAGATGGA         0,32         1         3,13         0,32         Hs.211577         kinectin 1 (kinesin receptor)           1495         TTTACCTTTGG         0,32         1         3,13         0,32         Hs.21108         ESTs           1496         TACGATATTCA         0,32         1         3,13         0,32         Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1498         GCTGCTAGAAA         0,32         1         3,13         0,32         Hs.197751         KIAA0666 protein           1499         GGGTAGAGAGT         0,32         1         3,13         0,32         Hs.196437         hypothetical protein FLJ10788           1500         ATCGGCTCCCA         0,32         1         3,13         0,32         Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501         ATAATGGAGTG         0,32         1         3,13         0,32         Hs.17850         ESTs           1503         TGCCTGACAAG         0,32         1         3,13         0,32         Hs.169160         ESTs           1504								
1494   AGAAAGATGGA	1402	TTTTOTOTATT	0.22		2.40	0.00	11- 04050	
1494 AGAAAGATGGA         0,32         1 3,13         0,32 Hs.211577         kinectin 1 (kinesin receptor)           1495 TTTACCTTTGG         0,32         1 3,13         0,32 Hs.21108         ESTs           1496 TACGATATTCA         0,32         1 3,13         0,32 Hs.207776         aspartylglucosaminidase           1497 GCACTGGGGCA         0,32         1 3,13         0,32 Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1498 GCTGCTAGAAA         0,32         1 3,13         0,32 Hs.197751         KIAA0666 protein hypothetical protein FLJ10788           1500 ATCGGCTCCCA         0,32         1 3,13         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32         1 3,13         0,32 Hs.17850         ESTs           1502 GAAGAAAGACT         0,32         1 3,13         0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955	1493	ITTICIGIATI	0,32	1	3,13	0,32	Hs.21356	
1495   TTTACCTTTGG   0,32	1404	ACAAACATOCA	0.00		0.40	0.00	11 044	
1495 TTTACCTTTGG         0,32         1 3,13         0,32 Hs.21108         ESTs           1496 TACGATATTCA         0,32         1 3,13         0,32 Hs.207776         aspartylglucosaminidase           1497 GCACTGGGGCA         0,32         1 3,13         0,32 Hs.206259         Homo sapiens mRNA for KlAA1190 protein, partial cds           1498 GCTGCTAGAAA         0,32         1 3,13         0,32 Hs.197751         KlAA0666 protein           1499 GGGTAGAGAGT         0,32         1 3,13         0,32 Hs.196437         hypothetical protein FLJ10788           1500 ATCGGCTCCCA         0,32         1 3,13         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32         1 3,13         0,32 Hs.17850         ESTs           1502 GAAGAAAGACT         0,32         1 3,13         0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955	1494	AGAAAGATGGA	0,32	7	3,13	0,32	Hs.2115//	·
1496 TACGATATTCA         0,32         1 3,13 0,32 Hs.207776         aspartylglucosaminidase           1497 GCACTGGGGCA         0,32         1 3,13 0,32 Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1498 GCTGCTAGAAA         0,32         1 3,13 0,32 Hs.197751         KIAA0666 protein           1499 GGGTAGAGAGT         0,32         1 3,13 0,32 Hs.196437         hypothetical protein FLJ10788           1500 ATCGGCTCCCA         0,32         1 3,13 0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32         1 3,13 0,32 Hs.17850         ESTs           1502 GAAGAAAGACT         0,32         1 3,13 0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13 0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13 0,32 Hs.169160         ESTs           1505 AAATATTAAAC         0,32         1 3,13 0,32 Hs.16364         hypothetical protein FLJ10955	1.105							
1497         GCACTGGGGCA         0,32         1         3,13         0,32         Hs.206259         Homo sapiens mRNA for KIAA1190 protein, partial cds           1498         GCTGCTAGAAA         0,32         1         3,13         0,32         Hs.197751         KIAA0666 protein hypothetical protein FLJ10788           1500         ATCGGCTCCCA         0,32         1         3,13         0,32         Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501         ATAATGGAGTG         0,32         1         3,13         0,32         Hs.17850         ESTs           1502         GAAGAAAGACT         0,32         1         3,13         0,32         Hs.172506         myosin VB           1503         TGCCTGACAAG         0,32         1         3,13         0,32         Hs.169160         ESTs           1504         TTCCTGTAATC         0,32         1         3,13         0,32         Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505         AAATATTAAAC         0,32         1         3,13         0,32         Hs.16364         hypothetical protein FLJ10955								
1498 GCTGCTAGAAA   0,32   1 3,13   0,32 Hs.197751   KIAA0666 protein								
1498 GCTGCTAGAAA         0,32         1 3,13         0,32 Hs.197751         KIAA0666 protein           1499 GGGTAGAGAGT         0,32         1 3,13         0,32 Hs.196437         hypothetical protein FLJ10788           1500 ATCGGCTCCCA         0,32         1 3,13         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32         1 3,13         0,32 Hs.17850         ESTs           1502 GAAGAAAGACT         0,32         1 3,13         0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955	1497	GCACTGGGGCA	0,32	1	3,13	0,32	Hs.206259	
1498 GCTGCTAGAAA         0,32         1 3,13         0,32 Hs.197751         KIAA0666 protein           1499 GGGTAGAGAGT         0,32         1 3,13         0,32 Hs.196437         hypothetical protein FLJ10788           1500 ATCGGCTCCCA         0,32         1 3,13         0,32 Hs.194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32         1 3,13         0,32 Hs.17850         ESTs           1502 GAAGAAAGACT         0,32         1 3,13         0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955			}				•	KIAA1190 protein, partial
1499 GGGTAGAGAGT         0,32         1 3,13 0,32 Hs.196437.         hypothetical protein FLJ10788           1500 ATCGGCTCCCA         0,32         1 3,13 0,32 Hs.194478.         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501 ATAATGGAGTG         0,32         1 3,13 0,32 Hs.17850.         ESTs           1502 GAAGAAAGACT         0,32         1 3,13 0,32 Hs.172506.         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13 0,32 Hs.169160.         ESTs           1504 TTCCTGTAATC         0,32         1 3,13 0,32 Hs.167106.         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13 0,32 Hs.16364.         hypothetical protein FLJ10955.							_	cds
1499         GGGTAGAGAGT         0,32         1         3,13         0,32         Hs. 196437         hypothetical protein FLJ10788           1500         ATCGGCTCCCA         0,32         1         3,13         0,32         Hs. 194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501         ATAATGGAGTG         0,32         1         3,13         0,32         Hs. 17850         ESTs           1502         GAAGAAAGACT         0,32         1         3,13         0,32         Hs. 172506         myosin VB           1503         TGCCTGACAAG         0,32         1         3,13         0,32         Hs. 169160         ESTs           1504         TTCCTGTAATC         0,32         1         3,13         0,32         Hs. 167106         proteasome (prosome, macropain) subunit, alpha type,           1505         AAATATTAAAC         0,32         1         3,13         0,32         Hs. 16364         hypothetical protein FLJ10955	1498	GCTGCTAGAAA	0,32	1	3,13	0,32	Hs.197751	KIAA0666 protein
The function of the function	1499	GGGTAGAGAGT	0,32	1	3,13	0,32	Hs.196437	·
1500         ATCGGCTCCCA         0,32         1         3,13         0,32         Hs. 194478         Homo sapiens mRNA; cDNA DKFZp434O1572 (from           1501         ATAATGGAGTG         0,32         1         3,13         0,32         Hs. 17850         ESTs           1502         GAAGAAAGACT         0,32         1         3,13         0,32         Hs. 172506         myosin VB           1503         TGCCTGACAAG         0,32         1         3,13         0,32         Hs. 169160         ESTs           1504         TTCCTGTAATC         0,32         1         3,13         0,32         Hs. 167106         proteasome (prosome, macropain) subunit, alpha type,           1505         AAATATTAAAC         0,32         1         3,13         0,32         Hs. 16364         hypothetical protein FLJ10955					,	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
CDNA DKFZp434O1572 (from   1501 ATAATGGAGTG   0,32	1500	ATCGGCTCCCA	0.32	1	3.13	0.32	Hs 194478	
1501 ATAATGGAGTG			-,	· l		-,		
1501 ATAATGGAGTG         0,32         1 3,13         0,32 Hs.17850         ESTs           1502 GAAGAAAGACT         0,32         1 3,13         0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955								
1502 GAAGAAAGACT         0,32         1 3,13         0,32 Hs.172506         myosin VB           1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955	1501	ATAATGGAGTG	0.32		3 13	0.32	Hs 17850	
1503 TGCCTGACAAG         0,32         1 3,13         0,32 Hs.169160         ESTs           1504 TTCCTGTAATC         0,32         1 3,13         0,32 Hs.167106         proteasome (prosome, macropain) subunit, alpha type,           1505 AAATATTAAAC         0,32         1 3,13         0,32 Hs.16364         hypothetical protein FLJ10955								<del> </del>
1504 TTCCTGTAATC 0,32 1 3,13 0,32 Hs.167106 proteasome (prosome, macropain) subunit, alpha type,  1505 AAATATTAAAC 0,32 1 3,13 0,32 Hs.16364 hypothetical protein FLJ10955								
macropain) subunit, alpha type,  1505 AAATATTAAAC 0,32 1 3,13 0,32 Hs.16364 hypothetical protein FLJ10955								
1505 AAATATTAAAC 0,32 1 3,13 0,32 Hs.16364 hypothetical protein FLJ10955	1304	I TOOTGIAATO	0,32	'	3,13	0,32	⊓s. 10/100	
1505 AAATATTAAAC 0,32 1 3,13 0,32 Hs.16364 hypothetical protein FLJ10955				ſ				
FLJ10955	4505	AAATATTAAAA	- 0.00		0.15	0.00	11 1000	
	1505	AAATATTAAAC	0,32	1	3,13	0,32	Hs.16364	
	455							
1506 ATAAAGCCGAA 0,32 1 3,13 0,32 Hs.159471 ZAP3 protein								17150
1507 AGGCTGAGGCG 0,32 1 3,13 0,32 Hs.156292 ESTs								

1508	TAGTGCTCTCA	.0,32	1	3,13	0,32	Hs.154424	deiodinase, iodothyronine, type II
1509	TTCATAAAAAA	0,32	1	3,13	0,32	Hs.154057	matrix metalloproteinase 19
1510	GTGGCTACAGT	0,32	1	3,13	0.32	Hs.151251	ESTs
	TAATCTTTCTT	0,32	1	3,13		Hs.151236	highly charged protein
	GATGGGGAAAT	0,32	1	3,13		Hs.14520	eukaryotic translation initiation factor 2C, 1
1513	ACTAAGTGCTA	0,32	1	3,13	0,32	Hs.132739	I-mfa domain-containing protein
1514	CGGTTATTTAA	0,32	1	3,13	0,32	Hs.119488	cystein-rich hydrophobic domain 2
1515	AGTGTGGGACT	0,32	1	3,13	0,32	Hs.118821	CGI-62 protein
1516	ATAGTTTAGCA	0,32	1	3,13	0,32	Hs.112157	ESTs
1517	CACCGAGACCA	0,32	1	3,13	0,32	Hs.107169	insulin-like growth factor binding protein 5
	AAATGACAATA	0,32		3,13		Hs.104904	ESTs
	TGACCAGGCGC	0,32	1	3,13	_	Hs.1019	parathyroid hormone receptor 1
1520	GGGCATCTCTT	12,19	38	3,12		Hs.76807	major histocompatibility complex, class II, DR alpha
1521	GTGGCGGGAGC	1,93	6	3,11	1,62	Hs.68257	general transcription factor IIF, polypeptide 1 (74k
1522	GAGAAACCCCG	12,89		3,10		Hs.5486	clone FLB5214
1523	AAATGCGAACA	1,29	4	3,10	1,16	Hs.5672	ESTs, Weakly similar to Similarity to Yeast D-lactat
1524	ATCCACCTGCC	0,97		3,09	0,91	Hs.231656	EST
1525	GCCAGCTGACA	0,97		3,09		Hs.118913	ESTs
1526	AGAAAGAATCT	0,97	3	3,09	0,91	Hs.107979	small membrane protein 1
1527	TGCCCCTTGCC	0,97	3	3,09		Hs.105700	secreted frizzled-related protein 4
	GAGAAACCCTG	11,34	35	3,09	7,42	Hs.194359	ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS
	TTGCTGACTTT	11,75		3,06		Hs.108885	collagen, type VI, alpha 1
1530	AGGCTGAGGCA	3,6	11	3,06		Hs.17834	downstream neighbor of SON
	TCACTGCATTC	1,64		3,05		Hs.235587	EST
	CTTAAAAAAAA	1,64		3,05		Hs.176626	hypothetical protein EDAG-1
	GTGAAAACCTG	2,96		3,04		Hs.283606	ESTs, Moderately similar to ALU5_HUMAN ALU
	AGCCACCGTGC	10,2		3,04		Hs.240845	DKFZP434D146 protein
1535	GTGGCACATAC	1,99	6	3,02	1,57	Hs.205353	ectonucleoside triphosphate diphosphohydrolase 1
1536	CCATTGTACTC	4,98	15	3,01	3.39	Hs.108740	DKFZP586A0522 protein
	TGCCTGTAATC	16,99		3,00		Hs.167135	Homo sapiens cDNA FLJ10728 fis, clone

## 288

4500	T	0.001	40	0.00	4 4 =	11- 044 453	IFOT
	TTAGCCAGGAT	6,33		3,00		Hs.211457	EST
	TAGGGAATGAA	0	3	3,00	4,35	Hs.59545	ring finger protein 15
1540	AAAGCATTTCT	이	3	3,00	4,35	Hs.36688	ESTs, Moderately similar
							to WAP four-disulfide core
1541	ATGACCCGCAG	0	3	3,00	4,35	Hs.286254	ESTs, Weakly similar to
							AF170723_1 protein
				<b> </b>			kinase ST
1542	ATTITITCAG	0	3	3,00	4,35	Hs.278004	EST
1543	GCAAGCCATTT	0	3	3,00	4,35	Hs.272813	dual oxidase 1
1544	GATTTTTTTT	0	3	3,00	4,35	Hs.227913	API5-like 1
	TCTCTTGGGGT	0	3	3,00	4.35	Hs.16740	hypothetical protein
				] -,]	.,		FLJ11036
1546	TGTGTGTAACA	0	3	3,00	4 35	Hs.156457	ESTs
	CCTTIGTCTTT	1	3	3,00	0.88	Hs.99654	protein-O-
1577	00111010111	'\	J	3,00	0,00	115.99004	mannosyltransferase 1
15/0	AATTGTAGTTA	1		3,00	0.00	He 6900	RAP2A, member of RAS
1048	MAIIGIAGIIA		3	3,00	0,08	Hs.6809	
4540	AGCCACTGTAC	<u> </u>		2.00	0.00	LIA EEO 40	oncogene family
1549	AGCCACTGTAC	1	3	3,00	0,88	Hs.55346	ESTs, Weakly similar to
							Z141_HUMAN ZINC
1070							FINGER
1550	CAAACTCAAAA	1	3	3,00	0,88	Hs.279809	hypothetical protein
		[					PRO1741
1551	GAGCACTTGGG	1	3	3,00	0,88	Hs.182937	peptidylprolyl isomerase A
							(cyclophilin A)
	AAGTTTTTAGT	1	3	3,00		Hs.149917	ESTs
1553	TTTGAGGATTG	1	3	3,00	0,88	Hs.147916	DEAD/H (Asp-Glu-Ala-
							Asp/His) box polypeptide 3
1554	TAGTTGGAACT	1	3	3,00	0,88	Hs.1119	nuclear receptor subfamily
					,		4, group A, member 1
1555	TTGACCAGGCT	4,34	13	3,00	2,98	Hs.285080	ESTs
	CTTATTTGTTT	1,67		2,99		Hs.4114	plastin 3 (T isoform)
	AGCTTCCAGCC	1,67	5	2,99		Hs.144974	ESTs, Highly similar to
7		.,	Ĭ	_,,,,,	.,		unnamed protein product
							IH.s
1558	CACCCCCTCGC	0,67	2	2,99	0.61	Hs.91246	hypothetical protein
,555		'',5'		2,00	0,01	10.01240	DKFZp547O146
1550	TCTCCAGGACA	0,67	2	2,99	0.61	Hs.8025	Homo sapiens clone
1009	·	0,07	2	2,33	0,01	1 13.0023	23767 and 23782 mRNA
1560	TATTTCAATTC	0.67		2 00	0.64	Hs.79507	sequences
	TATTTCAATTG	0,67		2,99			KIAA0582 protein
ומכו	CAGGTTGAAGT	0,67	2	2,99	0,61	Hs.79219	RalGDS-like gene;
4500	TATOTTAATOT	0.0=		0.55		70	KIAA0959 protein
1562	TATGTTAATGT	0,67	2	2,99	0,61	Hs.7341	ESTs, Weakly similar to
1==-						• • •	DUS8_HUMAN DUAL
	GACTGCTCTGG	0,67		2,99		Hs.36475	ESTs
	GAAGAGTGCTC	0,67	2	2,99		Hs.32204	ESTs
1565	GAGCCAAAGAA	0,67	2	2,99	0,61	Hs.29423	ESTs, Weakly similar to
			,				macrophage lectin 2
	<u> </u>		<u>L</u>				[H.sapie
1566	CTTGTAATCTC	0,67	2	2,99	0,61	Hs.278002	EST
						•	•

T 4 ===			<del></del>	0.55			1=0=
	TTACAATCACA	0,67		2,99		Hs.21276	ESTs
	GTGAAATCCAG	0,67		2,99		Hs.183275	ESTs
	CCTGTAATACC	3,02		2,98		Hs.92254	hypothetical protein FLJ20163
1570	AAAAGCAGAAA	1,35	4	2,96	1,11	Hs.84728	Kruppel-like factor 5 (intestinal)
1571	TTTGGGCCTAA	6,09		2,96		Hs.230320	EST
1572	CCTGTGGTCCC	17,64	52	2,95	10,04	Hs.249373	Homo sapiens clone FLB2543
	TAGCTCCCTTG	1,7		2,94		Hs.199160	myeloid/lymphoid or mixed-lineage leukemia (trithora
1574	GTGAGACCCTG	7,5		2,93	4,58	Hs.135756	polymerase (DNA- directed) kappa
1575	GCGAAACCCCG	22,18				Hs.30376	hypothetical protein
	CCACTGCATTC	12,98		2,93		Hs.270403	ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS
	GGGATCGCCCC	2,05		2,93		Hs.12865	p47
1578	TTCCAAGGCAG	3,08	9	2,92	2,13	Hs.317	topoisomerase (DNA) I
1579	ATGGTGGGTGC	3,08	9	2,92	2,13	Hs.209602	ESTs, Weakly similar to ubiquitous TPR motif, Y isof
1580	ATCTTGGCTCA	2,4	7	2,92	1,72	Hs.86958	interferon (alpha, beta and omega) receptor 2
1581	GTGAAACACCG	3,43		2,92	2,32	Hs.207766	EST
1582	CGTTCATTCAT	1,03	3	2,91	0,86	Hs.6139	synaptogyrin 1
1583	CCATAATGTTG	1,03	3	2,91	0,86	Hs.39957	pleckstrin 2 (mouse) homolog
1584	CTCTACGCATT	1,03	3	2,91	0,86	Hs.278573	H-2K binding factor-2
1585	ATGCAGAGGTG	1,03	3	2,91	0,86	Hs.210706	ESTs, Weakly similar to AF211175_1 unknown [H.sapien
1586	GCCAACAGCAT	1,03	3	2,91	0,86	Hs.155606	paired mesoderm homeo box 1
1587	CCTGTAATCAC	2,75	8	2,91	1,92	Hs.266136	ESTs
1588	GGATATGTGGT	7,24	21	2,90	4,34	Hs.738	early growth response 1
1589	GCTCACACCTG	1,38	4	2,90	1,09	Hs.60617	sialyltransferase 4A (beta- galactosidase alpha-2,3-s
1590	CATACAGAAAA	1,38	4	2,90		Hs.3107	CD97 antigen
1591	TATCCCAGAAC	9,32		2,90		Hs.169286	crystallin, beta B2
1592	CGCCTGTAATC	11,4		2,89		Hs.235083	EST
	GTGGCACACAC	16,96	49	2,89	9,23	Hs.71475	hypothetical protein
	ATCATACCACT	2,08	6	2,88	1,5	Hs.224071	EST, Weakly similar to ALUC_HUMAN IIII ALU CLASS C
	GTGGCGTGCAC	6,59		2,88		Hs.228163	EST
1596	TTGCCCAGGCT	17,35	50	2,88	9,38	Hs.56027	Homo sapiens mRNA; cDNA DKFZp586J1717 (from

4507	00007447000	0.40		0.00		11 070040	Internal constitution of a
	CCGGTAATCCC	2,43		2,88		Hs.272813	dual oxidase 1
	AGCCACTGTGC	8,35		2,87		Hs.180606	EST
1599	GTGGTGCACAC	16,79	48	2,86		Hs.272173	ESTs, Weakly similar to ALU1_HUMAN ALU
1600	CCCACTCTTTG	1,05	3	2,86		Hs.9414	KIAA1488 protein
1601	CAAAATCTTGA	1,05	3	2,86	0,84	Hs.75431	fibrinogen, gamma polypeptide
1602	CCTGTAGACCC	1,05	3	2,86	0,84	Hs.5123	inorganic pyrophosphatase
1603	TCCTGGTTATT	1,05	3	2,86	0.84	Hs.4084	KIAA1025 protein
	TGCTAGATTGG	1,05	3	2,86	0,84	Hs.239663	myeloid/lymphoid or mixed-lineage leukemia (trithora
1605	AACCCGGGAGA	1,05	3	2,86	0,84	Hs.236241	EST
	TACTCGGTTGT	1,05		2,86		Hs.119394	ESTs
	GACGGGGTGGA	1,05	3	2,86		Hs.111279	hypothetical protein
	ATTTGTGTGTA	0,7		2,86		Hs.94499	ESTs
	GTTCCAAGCAA	0,7		2,86		Hs.94011	ESTs, Weakly similar to MAGE-B4 [H.sapiens]
1610	CTATCTGTGGA	0,7	2	2,86	0,59	Hs.9176	ESTs
	GGCCCAGAGCC	0,7		2,86		Hs.91246	hypothetical protein DKFZp547O146
1612	TTGATGCCCAG	0,7	2	2,86	0,59	Hs.8503	ESTs
1613	TATTGTTAAAA	0,7	2	2,86	0,59	Hs.7984	ESTs
1614	CAATCTTTCAA	0,7		2,86		Hs.78909	butyrate response factor 2 (EGF-response factor 2)
1615	CTTCCTTGTGT	0,7	2	2,86	0,59	Hs.6298	KIAA1151 protein
	GACAGTGATAG	0,7	2	2,86		Hs.53913	hypothetical protein FLJ10252
1617	GGCCTCTGATG	0,7	2	2,86	0,59	Hs.46670	PRO1575 protein
	GCCTCCCCCAC	0,7	2	2,86	0,59	Hs.40109	KIAA0872 protein
	GGAGCAGACGC	0,7	2	2,86	0,59	Hs.31718	Homo sapiens cDNA FLJ11034 fis, clone PLACE1004258
1620	CTGCCCTCTGC	0,7	2	2,86	0.59	Hs.27801	zinc finger protein 278
	GGCTCTTCTGG	0,7	2	2,86	0,59	Hs.27721	hypothetical protein FLJ20353
1622	TTGCAATAGGT	0,7	2	2,86	0,59	Hs.25625	hypothetical protein FLJ11323
1623	TGATGATCATT	0,7	2	2,86		Hs.22394	hypothetical protein FLJ10893
	CCCAAACGGTA	0,7	2	2,86		Hs.195453	ribosomal protein S27 (metallopanstimulin 1)
	TTGGCCAAGAT	0,7		2,86		Hs.19522	hypothetical protein PRO2849
	TTTACCTGTTG	0,7	2	2,86	0,59	Hs.173381	dihydropyrimidinase-like 2
	TGTCAATGGGG	0,7		2,86		Hs.169055	golgi autoantigen, golgin subfamily a, 2
	CTTCCGGGTAA	0,7		2,86		Hs.108924	DKFZP586P1422 protein
1629	AAGGTTCTTCT	0,35	1	2,86	0,29	Hs.89695	insulin receptor

1000	TAATTITAAAA	0.051		2.00	0.00	Un 0004	TEST
	TAATTITAAAC	0,35		2,86		Hs.8861	ESTs
	TTAAATGCTCT	0,35	1			Hs.82501	similar to mouse Xrn1 / Dhm2 protein
1632	AGCTCCCAAGA	0,35	1	2,86	0,29	Hs.80475	polymerase (RNA) II (DNA directed) polypeptide J (13
1633	TGCTGCGGAAG	0,35	1	2,86	0,29	Hs.80306	Homo sapiens mRNA, clone:RES4-4
1634	GAGCAGTGCTG	0,35	1	2,86	0,29	Hs.7636	feline sarcoma (Snyder- Theilen) viral (v-fes)/Fujina
1635	CAAGGAAATGT	0,35	1	2,86	0,29	Hs.64840	ESTs
	GCAACACCGGA	0,35		2,86		Hs.63525	poly(rC)-binding protein 2
1637	GCCAAAGATGT	0,35	1	2,86	0,29	Hs.58636	squamous cell carcinoma antigen recognized by T cell
1638	GTTGCAGGCGC	0,35	1	2,86	0,29	Hs.5811	hypothetical protein FLJ20467
1639	TCTTTAAAAAA	0,35	1	2,86	0,29	Hs.55999	ESTs
-	TGTGCTTCTAG	0,35	1			Hs.38613	ESTs
J	TATCAAAAAA	0,35	1	2,86		Hs.32491	ESTs
	TAACTCCTAGT	0,35		2,86		Hs.31845	ESTs
	TGATATTAAAG	0,35	1			Hs.30661	electron-transferring- flavoprotein dehydrogenase
1644	AAACCAATTTT	0,35	1	2,86	0,29	Hs.30483	Homo sapiens mRNA; cDNA DKFZp434O1311 (from
1645	AAGAACTCAGG	0,35	1	2,86	0,29	Hs.30250	v-maf musculoaponeurotic fibrosarcoma (avian) oncoge
1646	CAGTCCCAAAA	0,35	1	2,86	0,29	Hs.29846	Human DNA sequence from clone 717M23 on
1647	AAACCGGTCCC	0,35	1	2,86	0,29	Hs.285490	ESTs, Weakly similar to unnamed protein product [H.s
1648	AACATTCCTAA	0,35	1	2,86	0,29	Hs.285429	ESTs
	TTAACATTTAT	0,35	1			Hs.279763	hypothetical protein FLJ10504
1650	ATGGCGTGTGC	0,35	1	2,86	0,29	Hs.278880	ESTs
-	GAGCTCTTCCT	0,35	1	2,86		Hs.274598	cytochrome P450, subfamily IID (debrisoquine, sparte
1652	GCAAGACCTTG	0,35		2,86		Hs.273603	ESTs
1653	AGGTTAAGAGA	0,35		2,86		Hs.272046	ESTs
	CCCGTAATCTC	0,35		2,86		Hs.270062	Homo sapiens mRNA; cDNA DKFZp586D0924 (from
1655	GCGGCACGCAC	0,35	1	2,86	0,29	Hs.269867	ESTs
	GACTTCCAGCA	0,35		2,86		Hs.265168	ESTs, Moderately similar to T10_MOUSE SER/THR-

1657	GCTGTTCTAAG	0,35	1	2,86	0,29	Hs.24422	regulatory factor X-
							associated protein
	GCACTGAGAAG_	0,35	1	2,86		Hs.239499	KIAA0185 protein
1659	GCGAGACCTTG	0,35		2,86		Hs.232157	ESTs
1660	TTGCATTCTCC	0,35		2,86		Hs.21379	ESTs
1661	TGCTTGTAGTC	0,35	1	2,86	0,29	Hs.209680	ESTs, Weakly similar to ALU1_HUMAN ALU
1662	AACGCAGCCTT	0,35	1	2,86	0,29	Hs.20060	KIAA0229 protein
1663	TAGGAAACCTG	0,35	1	2,86	0,29	Hs.200596	KIAA0547 gene product
1664	GGCTTTATTCT	0,35	1	2,86	0,29	Hs.200412	Homo sapiens mRNA; cDNA DKFZp434G0719 (from
1665	AATACTTCTCT	0,35	1	2,86	0,29	Hs.2003	T cell receptor beta locus
1666	GGTGACAGAGG	0,35	1	2,86	0,29	Hs.200235	ESTs
1667	AGAAAAAAAT	0,35	1	2,86		Hs.200057	ESTs, Weakly similar to ALU5_HUMAN ALU
1668	CATTGGCACTC	0,35	1	2,86	0,29	Hs.195614	splicing factor 3b, subunit 3, 130kD
1669	TTAATTAGCAA	0,35	. 1	2,86	0,29	Hs.183085	Homo sapiens mRNA; cDNA DKFZp434K098 (from clone
1670	CGCGTCAGAGC	0,35	1	2,86	0,29	Hs.182982	golgin-67
	AAAACTCGCCG	0,35		2,86		Hs.17969	KIAA0663 gene product
	TGTACTTTCCT	0,35		2,86		Hs.179661	tubulin, beta polypeptide
	CAGACTGGGAG	0,35		2,86		Hs.1790	nuclear receptor subfamily 3, group C, member 2
1674	TTTCTGAAGGG	0,35	1	2,86	0,29	Hs.172910	EST's, Highly similar to unnamed protein product [H.s
1675	TAAATAAGGAA	0,35	1	2,86	0,29	Hs.17235	ESTs
1676	GATGACAGAGT	0,35	1			Hs.171995	kallikrein 3, (prostate specific antigen)
1677	GTTACCGAGTG	0,35	1	2,86	0,29	Hs.171637	ESTs, Weakly similar to KIAA1317 protein [H.sapiens]
1678	GGTCAAATCAT	0,35	1	2,86	0,29	Hs.170162	KIAA1357 protein
1679	GCTTCACTTCC	0,35	1	2,86	0,29	Hs.164303	ESTs
	GACAATACACC	0,35	1	2,86	0,29	Hs.161554	hypothetical protein FLJ20159
1681	TTCCAAAAAA	0,35	1	2,86	0,29	Hs.159971	SWI/SNF related, matrix associated, actin dependent
1682	TGTGACCCCTC	0,35		2,86		Hs.159237	hexokinase 3 (white cell)
1683	TTTGTGCCATT	0,35	1	2,86		Hs.155507	ESTs
1684	CCTTGCCCTAT	0,35	1	2,86		Hs.143746	ESTs
1685	AGAACAAATAA	0,35	1	2,86		Hs.135721	ESTs
1686	AGACTGTACTG	0,35		2,86		Hs.132348	ESTs, Weakly similar to diaphanous 1 [H.sapiens]
1687	CCATCCCAGTG	0,35	1	2,86	0,29	Hs.127863	ESTs
	ACAGACTGTTA	0,35		2,86		Hs.125036	tumor endothelial marker 7

$\overline{}$		Γ					Inrocursor
1690	CCCCACCCATT	0.25	1	2.00	0.20	Up 40451	precursor
1009	GGCGACCCATT	0,35	1	2,86	0,29	Hs.12451	echinoderm microtubule-
4000	CACCTCCCATT	0.05	<del> </del> -	2.00	0.00	11: 44004	associated protein-like
1690	CAGGTCCCATT	0,35	1	2,86	0,29	Hs.11924	ESTs, Weakly similar to
1001			ļ				ALU1_HUMAN ALU
1691	ACAAAGAAAAG	0,35	1	2,86	0,29	Hs.118578	Homo sapiens cDNA
			ļ	ļ	ļ		FLJ20053 fis, clone
							COL00809
	ACTGATGCTCA	0,35		2,86		Hs.115467	ESTs
1693	TCCTCTTTCAA	0,35	1	2,86	0,29	Hs.113987	lectin, galactoside-binding,
	,						soluble, 2 (galectin 2)
1694	AGGCAGCACTG	0,35	1	2,86	0,29	Hs.11112	ESTs
1695	CTACTGCACTC	7,71	22	2,85	4,42	Hs.185989	ESTs
	CCCAGCTAATT	10,23	29	2,83		Hs.251235	EST
	GTGGCACGCAC	7,79		2,82		Hs.228343	EST
	GCAAAATCCCA	3,19		2,82		Hs.268051	ESTs, Weakly similar to
.000	00/00/00/00/0	0, 10	Ŭ	2,02	2.,04	113.200001	ALU2 HUMAN ALU
1600	GTGAAACCTTG	12,42	35	2,82	6.56	Hs.161554	hypothetical protein
1033	GIOMACCIIG	12,72	33	2,02	0,30	118.101334	FLJ20159
1700	ACTGTAATCCC	2 55	40	2 02	2.22	Ua 407900	ESTs .
		3,55		2,82		Hs.127809	<del></del>
	ATCGCACCACT	6,77		2,81		Hs.142569	ESTs
1702	GGGAAACCCCA	3,57	10	2,80	2,2	Hs.278281	ESTs, Weakly similar to
							alternatively spliced
							produc
	GTGAAACCCCT	9,64		2,80		Hs.229364	ESTs
	CAGCAGCAAAA	1,79		2,79		Hs.285090	ESTs
1705	TAGAAGCCAAC	2,52	7	2,78	1,63	Hs.7905	SH3 and PX domain-
	:						containing protein
				L			SH3PX1
1706	TTCTTTTTCTT	1,44	4	2,78	1,04	Hs.250722	(Manual assignment)
							MUG, Myeloid-upregulated
							protein
1707	CCTATAATCCT	1,44	4	2,78	1,04	Hs.158164	ATP-binding cassette,
		·		·			sub-family B (MDR/TAP),
		,					member
1708	TAAACGTGGCA	1,08	3	2,78	0.82	Hs.284146	hypothetical protein
		.,,		_,	1		DKFZp762N0610
1709	CAGAAGTCTTC	1,08	3	2,78	0.82	Hs.23921	ESTs, Weakly similar to
1,00	0,10,010110	1,00	Ŭ	_,.0	0,02	1.0.2002	ALU7 HUMAN ALU
1710	ACCAGCCAAAG	1,08	- 2	2,78	0.82	Hs.193090	ESTs, Highly similar to
'' '		1,00	ا ا	=, 10	0,02	1 13. 133030	AF161437_1 HSPC319
				1			[H.sapien
1711	GAAATGGGGAA	1,08		2 70	0.00	Hs.173933	
'''	OAN I GGGGAA	1,08	3	2,78	0,02	F18.17.3933	Homo sapiens mRNA for
			1			1	KIAA1439 protein, partial
1740	CTCTCCTATTO	4.00	<del></del>	0.70	0.00	Un 470440	cds
	GTGTGGTATTC	1,08	3	2,78	0,82	Hs.172140	ESTs
	CCGAGTTTTTG	1,08	3	2,78	0,82	Hs.139709	ESTs
	GGCAAACTTTA	1,08		2,78		Hs.102497	paxillin
1715	CCACAGGGGAT	5,42	15	2,77	3,05	Hs.119571	collagen, type III, alpha 1
	}	}	]	j			(Ehlers-Danios syndrome

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1716	CCTGTGGTCTC	2,17	6	2,76	1,44 	Hs.236504	EST, Weakly similar to ALU6_HUMAN ALU SUBFAMILY
1717	CACCACCACGC	1,82	5	2,75	1.23	Hs.5862	hypothetical protein
	GTGAAACCCGG	4,37		2,75		Hs.229170	ESTs
	GTGAAACCCTA	10,58		2,74		Hs.152081	ESTs
	ATATGTATATT	1,46		2,74	1.03	Hs.75839	zinc finger protein 6
							(CMPX1)
1/21	GTGAAACCACA	1,46	4	2,74	1,02	Hs.283788	hypothetical protein DKFZp547A023
1722	GGGATTAAAGC	1,46	4	2,74	1,02	Hs.211579	melanoma adhesion molecule
1723	TACCTTTGCTA	1,46	4	2,74	1,02	Hs.120980	nuclear receptor co- repressor 2
1724	AATGAATGAAA	0,73	2	2,74	0.57	Hs.8986	complement component 1,
					<u> </u>		q subcomponent, beta
1/25	ATATTTCATTC	0,73	2	2,74	0,57	Hs.79402	polymerase (RNA) II (DNA directed) polypeptide C (33
1726	TGGAGGGGCAG	0,73	2	2,74	0,57	Hs.7306	secreted frizzled-related protein 1
	TCGCGCAATAA	0,73	2	2,74	0,57	Hs.72249	protease-activated receptor 3
1728	CTCAAAATCAA	0,73	2	2,74	0,57	Hs.72165	hypothetical protein FLJ20283
1729	TGTACATATGT	0,73	2	2,74	0,57	Hs.268384	homolog of yeast CDH1/HCT1
1730	TGCAATGTTGT	0,73	2	2,74	0,57	Hs.171957	triple functional domain (PTPRF interacting)
1731	TCATTTTGTGA	0,73	2	2,74	0,57	Hs.154567	supervillin
1732	TAAAACGTGAA	0,73	2	2,74	0,57	Hs.12592	period (Drosophila) homolog 3
1733	TCATCTGCAAA	0,73	2	2,74	0,57	Hs.105189	ESTs, Weakly similar to AF148856_2 unknown [H.sapien
1734	GGCAAAATCTA	0,73	2	2,74	0,57	Hs.104627	Homo sapiens cDNA FLJ10158 fis, clone
1735	GACCTATCTCT	2,93	8	2,73	1,78	Hs.194431	palladin
1736	GTGGTGCGTGC	10,72	29	2,71	5,23	Hs.129727	X-ray repair complementing defective repair in Chine
	TCTTGAACAGC	1,11	3	2,70	0,8	Hs.72249	protease-activated receptor 3
1738	GGCTTTGGTCT	1,11	3	2,70	0,8	Hs.177592	ribosomal protein, large,
	CCACCACACCC	1,11	3	2,70	0,8	Hs.117582	CGI-43 protein
1740	ATGAAACCCCG	9,26		2,70		Hs.226396	hypothetical protein FLJ11126
1741	CCACTGCACTG	11,49	31	2,70	5.53	Hs.193220	ESTs
	CCACTGCGCTC	4,45		2,70			ESTs, Weakly similar to

		г					ALLIZ THISAASI ALLI
<del>                                     </del>	0.011.0000	14.15					ALU7_HUMAN ALU
1743	GAGAAACCCCA	11,13	30	2,70	5,36	Hs.5719	chromosome
					•		condensation-related
4= -				L	45 =		SMC-associated prote
	CCTGTAATCCT	23,38		2,69		Hs.165954	ESTs
1745	GACAGTCGGTG	1,49	4	2,68	1	Hs.8203	endomembrane protein
1=10		- 1					emp70 precursor isolog
	TTTTCTCTGAA	1,49		2,68		Hs.75516	tyrosine kinase 2
1747	TTGGCTAGGCC	2,61	7	2,68	1,57	Hs.211539	eukaryotic translation
1715							initiation factor 2, subunit
1748	CCCTTGTCCGA	2,61	7	2,68	1,57	Hs.127824	ESTs, Weakly similar to
							weak similarity to
1=15	10000100100						collagens
	AGCCCAGGAGT	3,37		2,67		Hs.274813	EST
	GTGGTGTGCAC	6,45		2,64		Hs.20126	KIAA0317 gene product
1751	CCTGTGATCCT	1,9	5	2,63	1,17	Hs.240395	potassium channel,
[							subfamily K, member 6
	0.070						(TWIK-2)
1752	CCTGTAAACCC	1,9	5	2,63	1,17	Hs.161554	hypothetical protein
				- 4-			FLJ20159
	TCAATAAAACC	1,52		2,63		Hs.151411	KIAA0916 protein
	ACGAAACCCCA	1,52		2,63		Hs.117582	CGI-43 protein
1755	TGACCACCCTT	1,14	3	2,63	0,78	Hs.42390	nasopharyngeal
ľ		Ì					carcinoma susceptibility
						<del></del>	protein
	CTCGAATAAAA	1,14	3	2,63		Hs.34871	KIAA0569 gene product
1757	CGACTGCACTC	1,14	3	2,63	0,78	Hs.182061	Novel human gene
							mapping to chomosome
						<del></del>	22
1758	CAGAATAATGT	1,14	3	2,63	0,78	Hs.125031	choline/ethanolaminephos
							photransferase
	CAGAAGGCCAC	0,76		2,63		Hs.8268	ESTs
1760	GAAAGAGCTCT	0,76	2	2,63	0,55	Hs.7337	hypothetical protein
							FLJ10936
1761	AAAATTATCTT	0,76	2	2,63	0,55	Hs.63657	hypothetical protein
					·		FLJ11005
1762	GGTGTCTGTGG	0,76	2	2,63	0,55	Hs.5889	ESTs, Weakly similar to
ļ		'					AC004876_5 similar to
							predic
1763	AACTGAGAAGT	0,76	2	2,63	0,55	Hs.56406	ESTs, Highly similar to
							unnamed protein product
1==							[H.s
	TGAGTGGTTTG	0,76		2,63		Hs.29672	ESTs
	GAAGTTGCCTT	0,76	2	2,63	0,55	Hs.26777	KIAA0843 protein
1766	TTGTTAAGCCT	0,76	2	2,63	0,55	Hs.26243	Homo sapiens cDNA
							FLJ11177 fis, clone
							PLACE1007402
	TATCTCAGAAC	0,76	2	2,63	0,55	Hs.223142	ESTs
	GGTGAATTTTA	0,76	2	2,63	0,55	Hs.210866	EST
1769	TGAGCACATAA	0,76	2	2,63	0,55	Hs.194208	suc1-associated

		<del></del> -					
							neurotrophic factor target 2 (FGFR s
	GTGCGTGCCTG	0,76	2	2,63	0,55	Hs.182354	ESTs
1771	ATTATCCAGCG	0,76	2	2,63	0,55	Hs.182225	RNA binding motif protein 3
	TCTTCTTTCAG	0,76		2,63	·	Hs.17757	Homo sapiens mRNA; cDNA DKFZp434E1515 (from
	CTCTCCAAACC	0,76		2,63		Hs.151242	complement component 1 inhibitor (angioedema, heredi
1774	CCATTGCTCTC	0,76	2	2,63		Hs.117582	CGI-43 protein
1775	AAGATCCTTGT	0,76	2	2,63	0,55	Hs.113503	karyopherin (importin) beta
1776	GGAACTTGGCT	0,76	2	2,63	0,55	Hs.105613	ESTs
	AGTTTGTCACC	0,76		2,63		Hs.10130	ESTs
1778	TCCACAGTGGG	0,38	1			Hs.99636	ESTs, Weakly similar to I54197 hypothetical protein
1779	TACCCCTCTCA	0,38	1	2,63	0,27	Hs.994	phospholipase C, beta 2
1780	AAAGATGTACA	0,38	1	2,63	0,27	Hs.95243	transcription elongation factor A (SII)-like 1
1781	ATTTATAATCC	0,38	1	2,63	0,27	Hs.914	major histocompatibility complex, class II, DP alpha
1782	ACTGTTTGTTT	0,38	1	2,63	0,27	Hs.814	major histocompatibility complex, class II, DP beta
	TCGATGTGGCG	0,38	- 1	2,63		Hs.81248	CUG triplet repeat, RNA- binding protein 1
	TTAAGATCTTC	0,38	1	2,63		Hs.79404	neuron-specific protein
	TTCTTGTCATA	0,38	1	2,63	0,27	Hs.79081	protein phosphatase 1, catalytic subunit, gamma isof
	AACAATTATCA	0,38	1	2,63		Hs.7845	Homo sapiens cDNA FLJ20820 fis, clone ADSE00490
	AGAAACACTCA	0,38	1	2,63	0,27	Hs.75782	general transcription factor IIIC, polypeptide 2 (be
1788	AGAAATAAAAA	0,38	1	2,63	0,27	Hs.74649	cytochrome c oxidase subunit VIc
	TAAGTGTGGTT	0,38	1	2,63	0,27	Hs.7327	claudin 1
1790	AGGAGAGAGCC	0,38	1	2,63		Hs.6932	Homo sapiens clone 23809 mRNA sequence
1791	TGGACAAGTCA	0,38	1	2,63	0,27	Hs.64988	ESTs
	TTAAACTGCTG	0,38	1	2,63		Hs.6232	KIAA0764 gene product
	AGACCTCACTG	0,38	1	2,63		Hs.49763	ESTs
	TATTTGTATTT	0,38	1	2,63		Hs.4764	KIAA0763 gene product
	TGGATAGATTC	0,38	1			Hs.45519	ESTs
	TTGATTGATTT	0,38	1	2,63		Hs.42927	Homo sapiens cDNA FLJ11298 fis, clone

·	Γ	т -					PLACE1009794
1707	ACTATATTGTG	0,38		2,63	0.27	Hs.42532	ESTs
	GAGTCCGGCCT	0,38	1			Hs.4069	glucocorticoid modulatory
				,			element binding protein 1
1799	TATTTATTTTT	0,38	1	2,63	0,27	Hs.39143	ESTs, Weakly similar to
					•		predicted using
				2 2 2			Genefinder [
	TGACATCCTGA	0,38		2,63		Hs.285056	ESTs
1801	CTGCAAGGACA	0,38	1	2,63		Hs.284135	Homo sapiens HSPC295 mRNA, partial cds
	ATCCCCCAGAA_	0,38		2,63		Hs.278386	ESTs
1803	CCACTGCGCTT	0,38	1	2,63	0,27	Hs.252836	EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY
1804	TGCCAGACCCT	0,38	1	2,63		Hs.249721	ESTs
1805	GGTGTGCACCT	0,38	1	2,63	0,27	Hs.24587	signal transduction protein (SH3 containing)
1806	CTTTATTTT	0,38	1	2,63	0,27	Hs.245710	heterogeneous nuclear ribonucleoprotein H1 (H)
1807	AGCGCTGGGGA	0,38	1	2,63		Hs.241471	RNB6
1808	CTAGGACCTGT	0,38	1	2,63	0,27	Hs.240112	KIAA0276 protein
1809	TAGTCCTAGCT	0,38	1	2,63	0,27	Hs.237372	EST
1810	ATTTAATTTTA	0,38	1	2,63	0,27	Hs.235883	ESTs
1811	GGCAACAAAGT	0,38	1	2,63	0,27	Hs.233364	ESTs
1812	CTGTAAGGATC	0,38	1	2,63	0,27	Hs.227730	integrin, alpha 6
	AGTCTTCCAGT	0,38	. 1	2,63	0,27	Hs.226213	cytochrome P450, 51 (lanosterol 14-alpha- demethylase
	GCTCCCCTCC	0,38		2,63		Hs.2157	Wiskott-Aldrich syndrome (ecezema- thrombocytopenia)
	AGTATTTATGA	0,38		2,63		Hs.203838	ESTs
1816	GCTAAACCCTG	0,38	1	2,63	0,27	Hs.202781	ESTs, Moderately similar to ALU5_HUMAN ALU
	CCAGCATTACC	0,38	1	2,00		Hs.20082	Homo sapiens zinc finger protein NY-REN-21 antigen m
	CCTGCAATCTC	0,38		2,63		Hs.197793	ESTs
	GAAAAATGCGC	0,38	1	2,63		Hs.193398	ESTs
	GCCAGGGCTCA	0,38	1			Hs.187913	ESTs, Moderately similar to MRP3 [H.sapiens]
	TAAAACTTACA	0,38	1	2,63	0,27	Hs.184075	ESTs
	ACCCTTTTTAT	0,38		2,63		Hs.183153	ADP-ribosylation factor 4- like
	GTTTCAAACGA	0,38		2,63		Hs.180535	ESTs, Weakly similar to S69890 mitogen inducible gen
1824	CAGTCTCAGTG	0,38	1	2,63	0,27	Hs.17767	Homo sapiens mRNA; cDNA DKFZp761N07121 (from c

1825	AAAAATTCATC	0,38	1	2,63		Hs.170328	moesin
1826	TAAATAAACAA	0,38	1	2,63	0,27	Hs.16755	MBIP protein
1827	CACCTCAAACA	0,38	1	2,63	0.27	Hs.157150	ESTs, Weakly similar to
					<b>'</b>		zinc finger protein 106
		1 1					IM.m
1828	AATGTCCTCGG	0,38	1	2,63	0.27	Hs.155987	KIAA0645 gene product
1829	TGTACCCCGCT	0,38	1			Hs.155975	protein tyrosine
		1		_,-,	-,		phosphatase, receptor
ł		1 1					type, C-assoc
1830	CGGGTTTGTGC	0,38	1	2,63	0.27	Hs.155482	hydroxyacyl glutathione
		-,		,	-,		hydrolase
1831	GTCCATCTTAA	0,38	1	2,63	0.27	Hs.153177	ribosomal protein S28
	TITATTTTTAG	0,38		2,63		Hs.152250	ESTs
	CCTAAAAAAAA	0,38	<u>_</u>	2,63		Hs.148907	Homo sapiens mRNA;
1000		0,50		2,00	0,21	119.140907	cDNA DKFZp564G223
							(from clone
1834	TGATCGAGCTT	0,38	1.	2,63	0.27	Hs.145867	ESTs
	GAAGATATTCC	0,38	1	2,63		Hs.14553	sterol O-acyltransferase
1033	GAAGAIATICC	0,30	1	2,03	0,21	ITS. 14000	(acyl-Coenzyme A:
	3						cholester
1926	ATGGAGCTGCA	0,38	1	2,63	0.27	Hs.142779	ESTs
	ATAAGACCTTA	0,38		2,63		Hs.142296	jerky (mouse) homolog
1838	CAAGCCAAAAA	0,38	7	2,63	0,27	Hs.14229	hypothetical protein
1000	04447000444			0.00		11 101000	FLJ10379
	CAAATGGCAAA	0,38		2,63		Hs.134292	ESTs
1840	GAAGTTTAAAT	0,38	1	2,63	0,27	Hs.132463	phosphoinositide-3-kinase,
1211							class 2, beta polypeptide
1841	GAAGGCAAGAT	0,38	1	2,63	0,27	Hs.1321	coagulation factor XII
							(Hageman factor)
	TTAGTTATGAC	0,38		2,63		Hs.13063	transcription factor CA150
	CTGTATGTTTA	0,38		2,63		Hs.128777	ESTs
1844	ACCTGCATTCC	0,38	1	2,63	0,27	Hs.125034	Homo sapiens cDNA
							FLJ10733 fis, clone
	CCAGCTGCCTG	0,38		2,63		Hs.11782	ESTs
1846	AAAAGTGGTGT	0,38		2,63		Hs.117582	CGI-43 protein
$\overline{}$	TTTATCTGATA	0,38		2,63		Hs.117582	CGI-43 protein
1848	TATTTTACCTA	0,38		2,63		Hs.114765	myeloid/lymphoid or
		1 1					mixed-lineage leukemia
							(trithora
1849	ACCTCCACACG	0,38	1	2,63	0,27	Hs.108947	KIAA0050 gene product
	GATTGCTGTGC	0,38		2,63		Hs.100555	DEAD/H (Asp-Glu-Ala-
					•		Asp/His) box polypeptide
			,				18 (Myc
1851	TGAAGCAGAAA	0,38	1	2,63	0,27	Hs.100407	Homo sapiens mRNA;
1							cDNA DKFZp564H2416
}							(from
1852	ATTGTGCCACT	10,64	28	2,63	4.88	Hs.272324	Homo sapiens mRNA;
		'-	•	'	,,	• • • • • • •	cDNA DKFZp566G231
							(from clone
1853	TCGGAGCTGTT	4,19	11	2,63	2.2	Hs.4055	chromosome 21 open
•	•		• •	. , 1	, -,-		· · · · · · · · · · · · · · · · · · ·

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<del>  1==</del>	0.0000000000000000000000000000000000000			0.55			reading frame 50
	GTGGTGTGCGC	3,43		2,62	·	Hs.278038	ESTs, Highly similar to PMM2_HUMAN
1855	TGCTACGAAAA	2,67	7	2,62	1,53	Hs.146550	myosin, heavy polypeptide
							9, non-muscle
1856	CCTGGCCTAAA	2,67	7	2,62	1,53	Hs.111676	protein kinase H11; small
							stress protein-like protei
1857	TTCACTGTGAG	21,42	56	2,61	8,99	Hs.621	lectin, galactoside-binding,
					•		soluble, 3 (galectin 3)
1858	CCCTACCCTGT	11,51	30	2,61		Hs.75736	apolipoprotein D
	GTGGCGTGCGC	2,31	6	2,60		Hs.117582	CGI-43 protein
1860	AGCCACCACAC	7,71	20	2,59	3,57	Hş.170310	cat eye syndrome
		4.	'				chromosome region,
	<u>:</u> .						candidate 1
1861	TCTACTAAAAA	1,93	5	2,59	1,15	Hs.48802	Homo sapiens clone
							23632 mRNA sequence
1862	GCCGGGCACGG	1,93	5	2,59	1,15	Hs.271480	hypothetical protein
							FLJ20686
	GTGCTCAAACC	1,93		2,59		Hs.103915	KIAA0346 protein
	CCACTGCACTT	13,54		2,58		Hs.194300	ESTs
1865	GTGGCGGACGC	1,55	4	2,58	0,96	Hs.182577	inositol polyphosphate-5-
1000		1.00					phosphatase, 75kD
1866	CCACTGCCCTC	4,66	12	2,58	2,31	Hs.1010	regulator of mitotic spindle
1007	07000070700	44.00			4.00		assembly 1
	GTGGCGTGTGC	11,69		2,57		Hs.278627	prenylcysteine lyase
1868	TCTGTAGTCCC	2,34	6	2,56	1,32	Hs.7358	Homo sapiens mRNA;
							cDNA DKFZp566D1146
1000	OTA ATTTA A OT	4.47		0.50	0.70	11 0404	(from
1869	CTAATTTAACT	1,17	3	2,56	0,76	Hs.9194	putative glialblastoma cell
1970	CTCTCACTCAT	4 4 7		0.50	0.70	11- 70040	differentiation-related
	GTCTCAGTCAT	1,17	3	2,56	0,76	Hs.78943	bleomycin hydrolase
10/1	GACAGTCACTC	1,17	3	2,56	0,76	Hs.6066	Rho guanine nucleotide
1972	CCTGTAGTCCA	1 17	2	2.56	0.76	Ha 277020	exchange factor (GEF) 4
		1,17	3	2,56 2,56	0,76	Hs.277028	EST
	CAGAGTTGTAT	1,17		2,50	0,76	Hs.109144	ESTs
	GGCTGAGCTCA GCTTTCTCAAA	1,96	5	2,55 2,55	1,13	Hs.83004	interleukin 14
<del></del>		1,96		_		Hs.177153	EST
10/0	CCTGTAGTCCT	7,85	20	2,55	3,48	Hs.179657	plasminogen activator,
1077	GTGAAACACTC	2 1 4		2 55	4.65	He 445057	urokinase receptor
10//	GTGAAACACTG	3,14	8	2,55	1,05	Hs.145357	ESTs, Moderately similar
1879	GTGAAGCCCCA	5,13	10	2 52	2 44	Hs.171501	to ALU7_HUMAN ALU
10/0	G I GAAGCCCCA	5,13	13	2,53	2,41	ms. 171501	ubiquitin specific protease
1870	CCTGTATTCCC	3 16	0	2 52	1 63	He 240749	ESTs
	TTCAGTGCCTG	3,16		2,53		Hs.249718	
	GGGAAACAGGT	2,37		2,53		Hs.180933	CpG binding protein
	CCTTTTTTTT	1,58		2,53		Hs.18368	DKFZP564B0769 protein
1002		0,79	2	2,53	0,53	Hs.9956	hypothetical protein
1882	GGGGCTTAGGA	0.70	2	2 52	0.50	Uc 90425	FLJ20259
	CTAGACAGTAA	0,79		2,53		Hs.89135	KIAA1528 protein
1 1004	IO I AGACAG I AA	0,79		2,53	<sub> </sub> ∪,ວ≾	Hs.52526	KIAA0669 gene product

1885	GTGTTCTGTGC	0,79	2	2,53	0.53	Hs.241567	RNA binding motif, single
1000		5,79		~,55	0,00	1.13.27 1007	stranded interacting prote
1886	TGCTGTAAAGG	0,79	2	2,53	0.53	Hs.23856	Homo sapiens HSPC091
		-,		_,	-,		mRNA, partial cds
1887	GAGGAGTGGGT	0,79	2	2,53		Hs.206770	zinc finger protein 297
1888	GTAAGACCCTG	0,79	2	2,53	0,53	Hs.164177	ESTs
1889	GGCCGTTAGAA	0,79	2	2,53	0,53	Hs.135	methylmalonate-
		i					semialdehyde
							dehydrogenase
	AGGCTAAAAGC	0,79		2,53		Hs.113029	ribosomal protein S25
	CTGTGTAATTT	0,79	2	2,53	0,53	Hs.109731	ESTs
1892	CTGAAGCGTGC	0,79	2	2,53	0,53	Hs.103391	Human insulin-like growth
							factor binding protein 5 (
1893	AGAACCTTCAA	3,57	9	2,52	1,78	Hs.181244	major histocompatibility complex, class I, A
1894	TTCTGTGCTGG	6,36	16	2,52	2,84	Hs.1279	complement component 1,
	-						r subcomponent
	TTAGCTGAGTC	1,99	5	2,51	1,12	Hs.153028	cytochrome b-561
1896	TTGGCCAGACT	3,6	9	2,50	1,76	Hs.91728	polymyositis/scleroderma
							autoantigen 1 (75kD)
1897	TTTCATTGCCT	3,6	9	2,50	1,76	Hs.173159	transforming, acidic coiled-
	1						coil containing protein
1898	GTGGCCAGAGG	3,6	9	2,50	1,76	Hs.1420	fibroblast growth factor
	•						receptor 3
4000	ACCOTTOTA			0.50	4.00	11 447500	(achondroplasia,
	ACCGTTCTGTA	2,4	- 6	2,50		Hs.117582	CGI-43 protein
	TAACTCCAAAG	1,2		2,50		Hs.24743	hypothetical protein FLJ20171
1901	TGCCGTAAATG	1,2	3	2,50	0,74	Hs.199067	v-erb-b2 avian
							erythroblastic leukemia
							viral oncogen
	TGAACTTTCCT	1,2	3	2,50	0,74	Hs.17567	ESTs
1903	TAAAGATCCTC	1,2	3	2,50	0,74	Hs.100407	Homo sapiens mRNA;
	<i>:</i>						cDNA DKFZp564H2416
1001	07000704040	00.40		0.40	2.44	11 100111	(from
1904	GTGGCTCACAC	22,18	55	2,48	8,11	Hs.138411	Homo sapiens mRNA;
				1			cDNA DKFZp586J1922
1905	GTGGTACACAC	2,02	E	2,48	11	Hs.250419	(from ESTs
	GATCTCTTGGG	2,02	5	2,48	1,1	Hs.115947	keratin 16 (focal non-
1900		2,02	. 5	2,40	1, 1	113.11394/	epidermolytic palmoplantar
·							ker
1907	GAGGAACTCAA	2,87	7	2,44	1.4	Hs.5008	CGI-87 protein
	AATAAAGCCTT	2,46		2,44		Hs.3314	selenoprotein P, plasma, 1
)	TTTACAAGTTA	1,64		2,44		Hs.91246	hypothetical protein
							DKFZp547O146
ļ	AGGTCAAAAAA	1,64		2,44		Hs.149570	actin related protein 2/3 complex, subunit 4 (20 kD)
1911	GAGCCCCCGTG	1,64	4	2,44	0,91	Hs.12908	CDC42-binding protein
l		l l		!			kinase beta (DMPK-like)

1012	GAGTAGCTGAG	1,23		2,44	0.72	Hs.260039	sarcospan (Kras
1912	GAGTAGCTGAG	1,23	3	<del>2,44</del>	0,72	HS.200039	oncogené-associated
ļ							gene)
1013	GTGCTGCTCCA	0,82	2	2,44	0.52	Hs.7936	BAI1-associated protein 2
	GAGATTTGTTT	0,82		2,44	0,52	Hs.75452	
				2,44	0,52		heat shock 70kD protein 2
	GAAGGGGTGCT	0,82	- 2	2,44	0,52	Hs.61950	DKFZp434A0131 protein
	GCCACAGTACA	0,82		2,44	0,52	Hs.55044	DKFZP586H2123 protein
	TAATTITTACT	0,82				Hs.52256	hypothetical protein FLJ20624
1918	ACTGTTTGGCA	0,82	2	2,44	0,52	Hs.286110	translocase of inner
		}	•				mitochondrial membrane 9
						· · · · · · · · · · · · · · · · · · ·	(yeast
	TCTGGCTAATT	0,82	2	2,44		Hs.262198	ESTs
1920	GTGGAAACCCA	0,82	2	2,44	0,52	Hs.243818	ESTs, Moderately similar
							to ALU1_HUMAN ALU
	TTGCCCAAGCT	0,82		2,44		Hs.213469	EST
	AAAACAGTGGC	0,82	2	2,44	0,52	Hs.184109	ribosomal protein L37a
1923	TGATGTGATAG	0,82	2	2,44	0,52	Hs.181159	Homo sapiens mRNA;
		ŀ			0		cDNA DKFZp434F0217
							(from
1924	TTTGAACCCTT	0,82	2	2,44	0,52	Hs.16206	uncharacterized
							hypothalamus protein
1000							HT008
1925	CCTATAATAAA	0,82	2	2,44	0,52	Hs.13885	ESTs, Weakly similar to
1000							T09A5.6 [C.elegans]
	AGAATCACTTA	0,82		2,44		Hs.130815	ESTs
	TATTTTGCAAA	0,82		2,44		Hs.11449	DKFZP564O123 protein
	CCTATAACCCC	0,41	1	2,44		Hs.99410	ESTs
1929	TCCAACTACAC	0,41	1	2,44	0,26	Hs.94581	sulfotransferase family,
4000	T4000444044	0.44			0.00	11 0100	cytosolic, 2B, member 1
1930	TACCCAAAGAA	0,41	1	2,44	0,26	Hs.9436	ESTs, Weakly similar to
1021	TOTTTOTAAAA	0.44		0.44	0.00	 Un 0074	NC5R_RAT NADH-5
	TGTTTGTAAAA	0,41		2,44		Hs.9271	KIAA1071 protein
1932	ттттттс	0,41	1	2,44	0,26	Hs.90797	Homo sapiens clone
4000	TATOTOTOGAA	- 44		0.44	0.00	11.00005	23620 mRNA sequence
	TATCTCTGCAA	0,41	1	2,44		Hs.82985	collagen, type V, alpha 2
	TTCTTCTGAAA	0,41		2,44		Hs.8087	NAG-5 protein
1935	CAGATGTTTAA	0,41	1	2,44	0,26	Hs.77631	glycine cleavage system
							protein H (aminomethyl
1026	TITGTAATATT	0.44	4	2 44	0.00	LID 7554C	carri
1930	TTTGTAATATT	0,41	1	2,44	0,20	Hs.75546	capping protein (actin
	1					•	filament) muscle Z-line,
1937	ACCCAGTTGTT	0,41	1	2,44	0.26	Hs.75410	heat shock 70kD protein 5
135/	, NOOMOTTOTT	0,71	'	۷,77	0,20	113.73410	(glucose-regulated protein
1938	CTCATTGGTGG	0,41	1	2,44	0.26	Hs.6580	Homo sapiens clone
.555	0/11/00/00	5,71	•	<u>ر</u> کر ج	0,20	13.0000	23718 mRNA sequence
1939	TCTTCTCACAA	0,41	1	2,44	0.26	Hs.656	cell division cycle 25C
	CCTTTGTTCAA	0,41		2,44		Hs.6107	ESTs 230
	TTAGAGATTCC	0,41		2,44		Hs.5947	mel transforming
, , , , , ,	1	0,71	•	, ~, <del>~</del>	5,20	113.0071	mor dansionning

		<del></del>				,	
							oncogene (derived from cell line NK
1942	TCCACACCAAA	0,41	1	2,44	0,26	Hs.53656	ESTs, Weakly similar to D29149 proline-rich protein
1943	TGTAATGGTTT	0,41	1	2,44	0,26	Hs.4930	low density lipoprotein receptor-related protein 4
1944	GTACTTACCTT	0,41	1	2,44	0,26	Hs.3454	ESTs, Weakly similar to KIAA0665 protein [H.sapiens]
1945	CTTAAATGGTT	0,41	1	2,44	0,26	Hs.29679	cofactor required for Sp1 transcriptional activation
1946	CTCCAACCTGA	0,41	1	2,44	0,26	Hs.285999	trinucleotide repeat containing 15
1947	TTAGGCTTTAG	0,41	1	2,44	0,26	Hs.285698	hypothetical protein FLJ20392
1948	GAAGATGTACG	0,41	1	2,44	0.26	Hs.285077	ESTs
	TGCCACCATAC	0,41		2,44		Hs.284138	ESTs
	TGTCTGTAGTC	0,41		2,44		Hs.282837	ESTs
	TAAAGTCCATT	0,41		2,44		Hs.278398	KIAA1117 protein
	CCAACTGACTT	0,41		2,44		Hs.277543	KIAA0631 protein
	ATGGAATGCTA	0,41	1	2,44		Hs.268551	receptor-interacting serine- threonine kinase 3
1954	TTGAAACCTCG	0,41	1	2,44	0.26	Hs.267148	ESTs
	TATATCATATT	0,41		2,44		Hs.266914	hypothetical protein FLJ10355
1956	TGGCACGCTGC	0,41	1	2,44	0,26	Hs.250890	ESTs, Weakly similar to TOM1 [H.sapiens]
1957	ACAGAGTCTCA	0,41	1	2,44	0,26	Hs.249031	EST
	GAAATATTGAT	0,41		2,44		Hs.247043	type 1 tumor necrosis factor receptor shedding amino
1959	GTGAAACCTGA	0,41	1	2,44	0,26	Hs.242076	EST
1960	ATGTCAACCAA	0,41	1	2,44		Hs.241558	ariadne (Drosophila) homolog 2
1961	GAAAAGGGCAC	0,41	1	2,44	0,26	Hs.23440	KIAA1105 protein
	TGCAGTCTTTG	0,41	1	2,44		Hs.232111	ESTs
	CAGATTTCCAG	0,41	1	2,44		Hs.21893	ESTs, Weakly similar to AF121081_1 cAMP inducible 2
1964	AAAAGGAAACC	0,41	1	2,44	0,26	Hs.21415	Homo sapiens mRNA; cDNA DKFZp761K2024 (from
1965	ATTGTAAGCTT	0,41	1	2,44	0,26	Hs.210232	ESTs
	GTCTTAAAATA	0,41		2,44		Hs.187991	DKFZP564A122 protein
	TCCGCAGGGAA	0,41	1	2,44		Hs.184592	Human clone A9A2BRB5 (CAC)n/(GTG)n repeat-
1968	GCTGGAGCTCA	0,41	1	2,44	0,26	Hs.181315	ESTs, Moderately similar to ALU4_HUMAN ALU
1969	TGTTGTAAATA	0,41	1	2,44	0,26	Hs.171501	ubiquitin specific protease

					Γ	l	111
1970	GCTCCTACATT	0,41	1	2,44	0.26	Hs.169488	dentatorubral-
1370		0,71	,	2,77	0,20	1115.105400	pallidoluysian atrophy
					}		(atrophin-1)
1971	CAGGTGCCAAA	0,41	1	2,44	0.26	Hs.168350	KIAA0554 protein
	TGCTGCTGCCC	0,41		2,44		Hs.167046	ESTs
	GCCTGGGCTGA	0,41	1			Hs.164476	hypothetical protein
							FLJ20626
	GTATGAGGTGG_	0,41		2,44		Hs.164464	ESTs
1975	GTAAACAĢAAA	0,41	1	2,44	0,26	Hs.161554	hypothetical protein FLJ20159
1976	TGAAATAAACT	0,41	1	2,44	0,26	Hs.155212	methylmalonyl Coenzyme
			_				A mutase
1977	TTTTGTCAACA	0,41	1	2,44	0,26	Hs.154645	ESTs, Weakly similar to
					•		tyrosine kinase [H.sapiens]
1978	GCACGTGTTCT	0,41	1	2,44	0,26	Hs.152096	cytochrome P450,
							subfamily IIJ (arachidonic
							acid epo
1979	TTAGTCCACAG	0,41	1	2,44	0,26	Hs.150390	zinc finger protein 262
1980	AAATTTCAAGC	0,41	1			Hs.146401	small inducible cytokine
	·				·		subfamily E, member 1
							(endo
1981	CCCAGCTACTT	0,41	1	2,44	0,26	Hs.143961	ESTs, Moderately similar
							to ALU1_HUMAN ALU
1982	AAACCAGGAAA	0,41	1	2,44	0,26	Hs.139120	ribonuclease P (30kD)
1983	AATTTCAAGAA	0,41	1	2,44	0,26	Hs.119591	adaptor-related protein
		·		·			complex 2, sigma 1
		·					subunit
1984	TACCCTAAAAT	0,41	1	2,44	0,26	Hs.117325	Homo sapiens cDNA
							FLJ11166 fis, clone
							PLACE1007242
1985	GGACTGTAGTG	0,41	1	2,44	0,26	Hs.11711	KIAA0329 gene product
1986	CTCCCGCCGGA	0,41	1	2,44	0,26	Hs.109445	KIAA1020 protein
1987	TGAAGTGCCCT	0,41	1	2,44	0,26	Hs.106932	ESTs
1988	GAAGAGGCTGG	0,41	1			Hs.105962	ESTs
1989	TTCCCTTCTTC	3,72	9			Hs.814	major histocompatibility
					,		complex, class II, DP beta
1990	AACCCGGAAGG	3,31	8	2,42	1,54	Hs.87497	butyrophilin, subfamily 3,
		'		'	,	•	member A2
1991	GTGGCGCGTGC	6,65	16	2,41	2.66	Hs.24135	hypothetical protein
		, ,					DKFZp761C241
1992	TACCCTAAAAC	42,92	103	2,40	13.65	Hs.165662	KIAA0675 gene product
	CCTGTGATCCC	16,67		2,40		Hs.249982	cathepsin B
	CCGGCCCTACC	1,67		2,40		Hs.271473	epithelial protein up-
		',-	٠	_, . •	] -,		regulated in carcinoma,
							membra
1995	GCACGCGTAAC	1,67	4	2,40	0.89	Hs.169552	ESTs, Weakly similar to
		''''		_, ,,	] 5,55		BRDT [H.sapiens]
1996	AACAAGGTGAG	1,26	3	2,38	0.71	Hs.94952	ESTs, Highly similar to
		',29		_,55	]	1.5.0 7002	transcription elongation fac
, _	1				1	ı	In mine a threat and in a digital and

1997	TGCGTCCCTCC	1,26	3	2,38	0,71	Hs.6179	DEAD/H (Asp-Glu-Ala- Asp/His) box polypeptide 17 (72k
1998	GACCCTTTGG	1,26		2,38		Hs.272848	Homo sapiens mRNA; cDNA DKFZp434G1310 (from
	GATTTTTCTGA	1,26	. 3	2,38		Hs.241567	RNA binding motif, single stranded interacting prote
	TGATTCTGTTT	1,26	_ 3	2,38	0,71	Hs.146428	collagen, type V, alpha 1
	CCCGGCTAATT	5,89		2,38		Hs.102926	ESTs
	GTGAAGCCCTG	6,74		2,37		Hs.105407	ectodermal dysplasia 1, anhidrotic
	ACTGAAAGAAG	2,11		2,37		Hs.169756	complement component 1, s subcomponent
	CTGAGAGCTGG	5,95		2,35		Hs.78501	growth arrest-specific 6
	GTTCCAGCAGC	1,7	4	2,35	0,88	Hs.23918	Homo sapiens clone 25116 mRNA sequence
2006	CCATTGCGCTC .	1,7		2,35		Hs.204299	ESTs, Moderately similar to alternatively spliced pr
	GGATGCGCAGG	1,7	4	2,35	0,88	Hs.168541	Homo sapiens mRNA full length insert cDNA clone EURO
	TTGGTGGAGGT	0,85		2,35		Hs.76294	CD63 antigen (melanoma 1 antigen)
	AGAATTATGGG	0,85		2,35		Hs.6975	PRO1073 protein
	TGTGGTGGCAC	0,85		2,35		Hs.46624	HSPC043 protein
	CAGTTCTTGAT	0,85	2	2,35	0,5	Hs.284217	serologically defined colon cancer antigen 33
2012	GCAAGACCCCG	0,85	2	2,35	0,5	Hs.262335	ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS
2013	AAAACAAAACA	0,85	2	2,35	0,5	Hs.24734	oxysterol binding protein
·	AAGACTGACAA	0,85	2	2,35	0,5	Hs.225951	topoisomerase-related function protein 4
	ттстсстсттт	0,85		2,35		Hs.22451	hypothetical protein FLJ10357
	TCAATCAGTGA	0,85		2,35		Hs.127270	ESTs
	TGGGGTCCCCA	0,85		2,35		Hs.123661	ESTs
	CCATTGCACTG	4,69		2,35		Hs.142457	ESTs, Moderately similar to alternatively spliced pr
2019	AATAAATTCCT	5,98	14	2,34	2,3	Hs.76307	neuroblastoma, suppression of tumorigenicity 1
	GTGGCAGATGC	2,99	7	2,34		Hs.273539	ESTs, Weakly similar to ALU2_HUMAN ALU
	CCTGTTATCCC	2,99	7	2,34		Hs.228142	EST
	GCGAAACTCCA	2,99		2,34		Hs.112860	KIAA1353 protein
	GTGAAACCTCG	16,26		2,34		Hs.194408	KIAA1244 protein
	GGCAGACACAT	1,29		2,33		Hs.33287	nuclear factor I/B
2025	AAAGAGAAGAG	1,29	3	2,33	0,69	Hs.22969	ESTs

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		1=1					
2026	ACAAAGCATTT	17,23	40	2,32	5,46	Hs.103391	Human insulin-like growth
							factor binding protein 5 (
2027	TTTAGTGACGT	3,02	7	2,32	1,32	Hs.7104	Kruppel-like factor 13
2028	AGCCACCACGC	7,35	17	2,31	2,63	Hs.60772	ESTs
2029	GCGAAACCTCG	3,46	8	2,31		Hs.210473	ESTs, Weakly similar to
		, , , ,		-,-			GELS_HUMAN
							GELSOLIN
2030	GCAGTTGGATC	1,73	1	2,31	0.86	Hs.284932	Homo sapiens clone
2000	COACTICOATO	1,73	7	2,01	0,00	113.204302	24650 ubiquitin hydrolase
Į.							mRNA, p
2024	CTTCTCAACTC	4 70		2.24	0.00	Hs.283681	
	CTTGTGAAGTG	1,73		2,31			ESTs
	GTGGCTCACGC	24,38		2,30		Hs.228230	EST
2033	CAGCTATTTCA	4,39	10	2,28	1,69	Hs.153179	fatty acid binding protein 5
							(psoriasis-associated)
2034	CACACACACAC	1,76	4	2,27	0,84	Hs.63984	cadherin 13, H-cadherin
							(heart)
2035	CCACCACACTC	1,76	4	2,27	0.84	Hs.256696	EST, Weakly similar to
		','	·	-,			ALU1_HUMAN ALU
		l l					SUBFAMILY
2036	CAAGGGTGACA	1,76		2,27	0.84	Hs.170222	solute carrier family 9
2030	CANGGGTGACA	1,70	-7	2,21.	0,04	1 15. 17 0222	
							(sodium/hydrogen
0007	COACCATAT	0.00		0.07	0.40	11. 00005	exchanger),
	GCAGCACTTAT	0,88	2	2,27		Hs.82035	GAP-like protein
2038	CCTGACCTCAA	0,88	2	2,27	0,48	Hs.7874	muskelin 1, intracellular
							mediator containing kelch
2039	GGGGTATGGTT	0,88	2	2,27	0,48	Hs.76144	platelet-derived growth
							factor receptor, beta
							polype
2040	AGCAGCCGCTC	0,88	2	2,27	0,48	Hs.7104	Kruppel-like factor 13
	CTCCTGGCCCA	0,88	2	2,27		Hs.5321	ARP3 (actin-related
		5,55		_,	0, .0		protein 3, yeast) homolog
2042	TTATGCCTCCA	0,88	2	2,27	0.48	Hs.43314	ESTs
	TACTTCCTGCG	0,88		2,27		Hs.38039	ESTs
			2				
2044	CCTTGCCCAGG	0,88	2	2,27	0,48	Hs.3144	Cas-Br-M (murine)
		<b> </b>				,	ectropic retroviral
							transforming s
2045	GTAGGGTTCCT	0,88	. 2	2,27	0,48	Hs.278597	protein tyrosine
	ļ	l i		[			phosphatase, non-receptor
				ļ		•	type 18 (
2046	ATAACCAAATG	0,88	2	2,27	0.48	Hs.25726	transposon-derived
				1	,		Buster1 transposase-like
	·	, ,					protein
2047	CCTGTGAATAG	0,88	2	2,27	0.48	Hs.200647	EST
	TGACCAGGGTC	0,88		2,27		Hs.167827	Homo sapiens clone
2070	INDROGRAGA	0,00	2	~,~ 1	U, <del>-1</del> 0	10.107.027	HH419 unknown mRNA
2040	CAGACCCAAAA	0.00		2 27	0.40	Ho 167550	·
	CAGACCCAAAA	0,88		2,27	0,48	Hs.167558	zinc finger protein 161
2050	TTCTCATAATC	0,88	2	2,27	J U,48	Hs.164919	ESTs, Highly similar to
							KPC2_HUMAN PROTEIN
							,
2051	ACTGATGCAAG	0,88	2	2,27	0,48	Hs.161049	ESTs
		-					

		2 2 2					E
2052	AACTCTGATAT	0,88	2	2,27	0,48	Hs.151046	hypothetical protein FLJ11193
2053	GGGCATCTCCA	0,88	2	2,27	0,48	Hs.107000	ESTs
2054	TGGATGTCTGT	0,44	1	2,27	0,24	Hs.96716	ESTs
2055	TATTAGAAGCA	0,44	1	2,27	0,24	Hs.91065	hypothetical protein DKFZp761B2423
2056	CACTGAATATG	0,44	1	2,27		Hs.86948	small nuclear ribonucleoprotein D1 polypeptide (16kD
2057	AGTTAATAAAG	0,44	1	2,27		Hs.8065	Homo sapiens mRNA full length insert cDNA clone EURO
2058	TTATATTTTCT	0,44	1	2,27	0,24	Hs.8021	KIAA1058 protein
2059	GTCACCAAACA	0,44	1	2,27	0,24	Hs.79283	selectin P ligand
2060	TTACACTGTAA	0,44	1	2,27	·	Hs.78687	neutral sphingomyelinase (N-SMase) activation associ
	AAGTTGCATCT	0,44	1	,		Hs.74649	cytochrome c oxidase subunit VIc
	TAGACATTTGA	0,44	1	2,27	0,24	Hs.74649	cytochrome c oxidase subunit VIc
	TTCATAGGTAA	0,44	1	2,27		Hs.7312_	ESTs
2064	CATTITAGGCA	0,44	. 1	2,27	0,24	Hs.72782	hypothetical protein FLJ11171
2065	ATACTATAATT	0,44	1	2,27		Hs.6966	Human DNA sequence from clone RP1-187J11 on
2066	CTTCGCTTTGT	0,44	1	2,27	0,24	Hs.69485	ESTs, Weakly similar to similar to other protein pho
2067	ATGATATATGC	0,44	1	2,27	0,24	Hs.6831	Homo sapiens clone 1400 unknown protein mRNA, partia
2068	GCCCCCTTACA	0,44	1	2,27	0,24	Hs.49476	Homo sapiens clone TUA8 Cri-du-chat region mRNA
2069	TCAGTCCCTGT	0,44	1	2,27	0,24	Hs.3685	hypothetical protein FLJ20209
	AATTITCATTA	0,44	1	2,27		Hs.35092	ESTs
	GCCAGTCCACT	0,44	1	2,27	0,24	Hs.34782	ESTs, Moderately similar to transducin [H.sapiens]
	TAGAGAGTTTA	0,44		2,27	0,24	Hs.29643	ESTs
	GCTGTCCCCTC	0,44	1			Hs.278422	DKFZP586G1122 protein
	GTGAAAGCCGT	0,44	1	2,27	0,24	Hs.270662	ESTs, Weakly similar to transformation-related prote
2075	TCTTCCAGAAA	0,44	1		0,24	Hs.256585	ESTs
2076	TTGGCCGGGAT	0,44	1			Hs.254900	ESTs
	GTAAAGAATGT	0,44	1	2,27	0,24	Hs.24790	ESTs
2078	AAATTTTTGTA	0,44		2,27		Hs.24650	ESTs, Moderately similar to AF133913_1 ARL-6 interac
2079	AACGCTGCAAA	0,44	1	2,27	0,24	Hs.24174	KIAA0876 protein

2000	CTACCCTAAAA	0.44	4	0.07	0.04	11- 000070	IECTa Manleha aimilea ta
2080	GTACCCTAAAA	0,44	1	2,27	0,24	Hs.239970	ESTs, Weakly similar to b34l8.1 [H.sapiens]
2081	TAATCTTTTTT	0,44	1	2,27	0.24	Hs.231463	EST
	AAATTGTATGT	0,44				Hs.22826	tropomodulin 3
2002	MAIIGIAIGI	0,77	'	2,21	0,24	115.22020	(ubiquitous)
2083	CCACTACATTC	0,44	1	2,27	0.24	Hs.22573	ESTs
	GCCGCACTCAG	0,44				Hs.200577	ESTs
	TCTTGTCATAC	0,44		-,-:		Hs.198998	conserved helix-loop-helix
2005	ICHGICAIAC	0,44	'	2,21	0,24	ns. 190990	lubiquitous kinase
2086	CAGCACCTGAT	0,44	1	2,27	0,24	Hs.198281	pyruvate kinase, muscle
2087	TTGAATAAAAG	0,44	1	2,27	0,24	Hs.198161	phospholipase A2, group IVB (cytosolic)
2088	GGGAAGTGTGC	0,44	1	2,27	0,24	Hs.197733	ESTs, Weakly similar to TRP7_HUMAN TRANSIENT
2089	CCTGGCCTACC	0,44	1	2,27	0,24	Hs.19585	KRAB-zinc finger protein SZF1-1
2090	TITATITTCAA	0,44	1	2,27	0,24	Hs.194293	ESTs
	TCTCTCTGCCT	0,44		2,27		Hs.184987	ESTs
2092	AGAGGAAGTAA	0,44				Hs.177537	ESTs, Weakly similar to ALU1_HUMAN ALU
2093	ACCAAAAAAA	0,44	1	2,27	0,24	Hs.173724	creatine kinase, brain
	GGGACATTTAT	0,44				Hs.173108	Homo sapiens clone 24523 mRNA sequence
2095	TACCATCCATA	0,44	1	2,27	0,24	Hs.169476	glyceraldehyde-3- phosphate dehydrogenase
2096	TTGATGAAGAA	0,44	1	2,27	0,24	Hs.168075	karyopherin (importin) beta 2
2097	GCCCGGTGCCC	0,44	1	2,27	0,24	Hs.1665	zinc finger protein homologous to Zfp-36 in mouse
2098	AAGAACTGTTT	0,44	1	2,27	0,24	Hs.159456	zinc finger protein 288
2099	CCAGCAACTGT	0,44			0,24	Hs.145279	SET translocation (myeloid leukemia-associated)
2100	ATGGTGCGTGC	0,44		2,27		Hs.140498	ESTs, Weakly similar to ALUE_HUMAN !!!! ALU CLASS
	CTCTTCAGGGT	0,44		2,27		Hs.13781	Homo sapiens cDNA FLJ11302 fis, clone PLACE1009971
2102	TCAATGTGAAA	0,44	1	2,27	0,24	Hs.13467	Homo sapiens BAC clone RP11-121A8 from 7p14- p13
2103	CTTAATACTAC	0,44	1	2,27	0,24	Hs.13273	KIAA0592 protein
	GGCTGCAGTAT	0,44				Hs.129892	KIAA0522 protein
	GGAAGCTGAAG	0,44				Hs.128629	ESTs
	CCTCGGGCATC	0,44		2,27		Hs.126735	ESTs
	TTTTCTTGCTG	0,44		2,27		Hs.120907	Homo sapiens mRNA; cDNA DKFZp547D135 (from clone

2400	CCCTTCCCTCT	0.44	4	2 27	0.24	Un 110027	ESTs
	GCCTTGCCTCT	0,44		2,27		Hs.118837	
	CTTTAAAAAAA	0,44		2,27		Hs.118162	fibronectin 1
	TGCGGAAAAA	0,44		2,27		Hs.113207	G protein-coupled receptor 30
2111	TGACTTTCTGC	0,44	1	2,27	0,24	Hs.11123	ESTs, Weakly similar to B38919 hypothetical protein
	CATTTACATAT	0,44		2,27		Hs.109438	Homo sapiens clone 24775 mRNA sequence
2113	AATGTGTTACT	0,44	. 1	2,27	0,24	Hs.105751	Ste20-related serine/threonine kinase
2114	ATCTTTATTCC	0,44	1	2,27	0,24	Hs.10351	KIAA0308 protein
	TGTTTGAATTC	0,44		2,27		Hs.103422	Homo sapiens mRNA; cDNA DKFZp434F1622 (from
	ACAACACCCCA	2,2	5	2,27		Hs.21453	Homo sapiens mRNA for inositol 1,4,5-trisphosphate 3
	CTGGAAATAAA	1,32		2,27		Hs.69745	ferredoxin reductase
	TTATTTATGAA	1,32	3	2,27		Hs.245188	tissue inhibitor of metalloproteinase 3 (Sorsby fund
	TACCAAGGATT	1,32	3	2,27		Hs.21729	splicing factor 3a, subunit 1, 120kD
2120	GTGCCAAACAC	1,32	1	2,27		Hs.172216	chromogranin A (parathyroid secretory protein 1)
	CCCGGCCCAAA	1,32	3	2,27	0,67	Hs.133207	PTPRF interacting protein, binding protein 1 (liprin
2122	AGAATTGCTTG	16,35	37	2,26 	4,88	Hs.56542	X-prolyl aminopeptidase (aminopeptidase P) 1, solubl
2123	TCTCTGATGCT	15;06	34	2,26	4,52	Hs.6441	tissue inhibitor of metalloproteinase 2
2124	ACACTGCACTC	3,11	7	2,25	1,27	Hs.200454	ESTs, Weakly similar to ALU1_HUMAN ALU
	GAAATGAGCAG	2,67	6	2,25	1,13	Hs.77293	KIAA0127 gene product
	GCTGGATGCGG	2,67	6	2,25		Hs.18075	chromosome 9 open reading frame 3
2127	GCAAAACCCTA	2,67		2,25		Hs.108740	DKFZP586A0522 protein
2128	TTTGCTCTCCC	6,68		2,25		Hs.75350	vinculin
	GGAGGCTGAGG	21,42		2,24		Hs.185973	membrane fatty acid (lipid) desaturase
	GTAAAACCCCA	9,38		2,24		Hs.18955	Homo sapiens cDNA FLJ20667 fis, clone KAIA596
	GGAGGGGCTT	13,42	30	2,24		Hs.77886	lamin A/C
2132	AGCTAAGTTTG	1,79	4	2,23	0,83	Hs.19447	ESTs, Weakly similar to CL36_HUMAN LIM DOMAIN

2422	CACACAGTTTT	0.06	10	2 22	2.62	H- 204254	rea homeles cono family
		8,06		2,23		Hs.204354	ras homolog gene family, member B
	ATGGCGGGTGC	4,04		2,23		Hs.172382	hypothetical protein FLJ20001
	GACTTGTATAT	2,7		2,22	·	Hs.81328	nuclear factor of kappa light polypeptide gene enhan
	ATAGTACAGCC	1,35		2,22		Hs.6361	MEK partner 1
	AGCCACCGCTC	1,35		2,22		Hs.6195	ESTs
	GGCAGGATGAT	1,35		2,22		Hs.274319	hypothetical protein FLJ10509
2139	GCCTGGGACCT	1,35	3	2,22	0,66	Hs.180871	protein kinase C, alpha binding protein
2140	ACAGCCGTGGG	1,35	3	2,22	0,66	Hs.123090	SWI/SNF related, matrix associated, actin dependent
	AGCCGAGATCA	2,26		2,21		Hs.277663	EST, Weakly similar to ALU1_HUMAN ALU SUBFAMILY
	AGCCTTTGTTG	11,78	26	2,21		Hs.9930	collagen-binding protein 2 (colligen 2)
2143	GTGTGCCTCCA	2,72	6	2,21	1,1	Hs.75254	interferon regulatory factor 3
2144	TGGAGAAGAGC	3,63	8	2,20	1,37	Hs.179526	upregulated by 1,25- dihydroxyvitamin D-3
2145	GGCCCCATTGC	1,82	4	2,20	0,81	Hs.173421	Human clone CE29 8.1 (CAC)n/(GTG)n repeat- containing
·2146	AATATTCATAG	0,91	2	2,20	0,47	Hs.8583	similar to APOBEC1
2147	GCTCCGTAAGG	0,91	2	2,20	0,47	Hs.80712	KIAA0202 protein
2148	GAGGTGCCCCA	0,91	2	2,20	0,47	Hs.77955	ESTs
2149	AGTGTATTTT	0,91	2	2,20	0,47	Hs.76473	insulin-like growth factor 2 receptor
2150	TAAACCTAAAG	0,91	2	2,20	0,47	Hs.60548	hypothetical protein PRO1635
2151	AGTCAAGCCCC	0,91	2	2,20	0,47	Hs.57687	four and a half LIM domains 3
	TATCAAAACAT	0,91	2	2,20	0,47	Hs.258939	EST
	ATCCTACTGTT	0,91	2	2,20	0,47	Hs.239218	uncharacterized hypothalamus protein HCDASE
	GCACCAAATGA	0,91		2,20		Hs.23585	KIAA1078 protein
2155	CCAATGCTATG	0,91	2	2,20	0,47	Hs.22753	Homo sapiens mRNA; cDNA DKFZp434K0926 (from
	CATCGTTACAT	0,91		2,20		Hs.173802	KIAA0603 gene product
2157	TAACCAAAAAC	0,91	2	2,20	0,47	Hs.169241	ELK4, ETS-domain protein (SRF accessory protein 1)
2158	CTGAAACCCCA	0,91	2	2,20	0,47	Hs.162132	ESTs
	GGAAGGCAAGC	0,91		2,20		Hs.144998	ESTs, Weakly similar to

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2460	CAAACCTCCTT	0.04		2.20	6.47	Up 44204	ALU7_HUMAN ALU
	GAAAGGTGGTT	0,91		2,20	<u> </u>	Hs.14394	hypothetical protein FLJ20157
2161	TCATAACCTTG	0,91	. 2	2,20	0,47	Hs.124029	inositol polyphosphate-5- phosphatase, 40kD
2162	TACCCCATAAA	2,29	5	2,18	0,96	Hs.281083	ESTs
	CGCCCCTGCG	2,75	6	2,18	1,09	Hs.135805	ESTs, Weakly similar to KIAA1323 protein [H.sapiens]
2164	ACAAAGCCCCA	1,38	3	2,17 2,17	0,65	Hs.8583	similar to APOBEC1
	CCTATAGTCCT	1,38				Hs.41694	origin recognition complex, subunit 2 (yeast homolog
	CCAGTACAGCC	1,38		2,17		Hs.140978	Homo sapiens mRNA; cDNA DKFZp762H106 (from clone
	CTGCTGCTGGT	1,38		2,17		Hs.12289	Cdc42 effector protein 2
	CTTAATCTTGT	1,85		2,16		Hs.75462	BTG family, member 2
	GCCACACCCCC	1,85		2,16		Hs.113916	Burkitt lymphoma receptor 1, GTP-binding protein
	TCCTCCCTACT	6,48		2,16		Hs.70266	yeast Sec31p homolog
	GTGAGACCCCA	5,57	12	2,15	1,8	Hs.198671	ESTs
	CAGATGCAAAA	5,13	0	2,14		Hs.89506	paired box gene 6 (aniridia, keratitis)
2173	CCCCAGGAGAA	2,34	5	2,14	0,93	Hs.169902	solute carrier family 2 (facilitated glucose transpo
2174	CTTCTGGGGAC	2,81	6	2,14	1,06	Hs.75082	ras homolog gene family, member G (rho G)
2175	GTGGCTTACAC	1,88	4	2,13	0,79	Hs.264482	Homo sapiens mRNA; cDNA DKFZp761A0411 (from
2176	GCGAGACCCCA	1,88	4	2,13	0,79	Hs.15681	ESTs
2177	TAAACTATTGG	1,41		2,13		Hs.78851	KIAA0217 protein
2178	GACGGCTGCAA	1,41		2,13		Hs.4909	dickkopf (Xenopus laevis) homolog 3
	GTGAAACCGTC	1,41		2,13		Hs.30596	Homo sapiens mRNA full length insert cDNA clone EURO
2180	CCACTGGACTC	1,41	3	2,13	0,63	Hs.253913	ESTs, Weakly similar to ALU1_HUMAN ALU
2181	TGTTAGCAAAT	1,41		2,13		Hs.22666	ESTs
	AAGTACGAGGA	1,41	3	2,13	0,63	Hs.22660	ESTs
	CTGTTGGAAAA	1,41		2,13		Hs.209863	ESTs
2184	AACCACTGTGC	1,41		2,13		Hs.188037	ESTs, Moderately similar to ALU7_HUMAN ALU
	CTTCTCCAAAA	0,94		2,13		Hs.99949	prolactin-induced protein
	GTGGTCAAGTT	0,94	2	2,13	0,46	Hs.92127	ESTs
L	CCTTTGCTGAG	0,94	2	2,13	0,46	Hs.7442	Human DNA sequence from clone 742C19 on
2188	TTGCTGCCAGC	0,94	2	2,13	0,46	Hs.5566	gap junction protein, beta 2, 26kD (connexin 26)

	TGAATTTCCTG	0,94	2	2,13	0,46	Hs.30057	Homo sapiens clone 24749 and 24750 mRNA sequences
	GTGGTATATGC	0,94	_	2,13		Hs.279893	hypothetical protein FLJ20342
	GTGGTGCACTC	0,94		2,13		Hs.268573	Homo sapiens mRNA; cDNA DKFZp762N226 (from clone
	GTGGCACAAGC	0,94	2	2,13	0,46	Hs.258487	EST
	AGCCACCATAC	0,94	2	2,13	0,46	Hs.236051	EST
	TAAAACCGTTT	0,94		2,13		Hs.182280	MADS box transcription enhancer factor 2, polypeptid
2195	CCTTGTTTAAC	0,94	. 2	2,13	0,46	Hs.173965	ribosomal protein S6 kinase, 90kD, polypeptide 3
2196	GCTCACTGCAA	0,94	2	2,13	0,46	Hs.163385	EST
	TGCCTTGGGCT	0,94		2,13 2,13		Hs.1497	retinoic acid receptor, gamma
	TTATTGTTCCC	0,94		2,13		Hs.12126	hepatocellular carcinoma- associated antigen 112
	GGGACAACCCA	0,94	2	2,13	0,46	Hs.11530	ESTs
	GTTTCTTACTG	0,94	2	2,13	0,46	Hs.106204	KIAA1327 protein
	TTTTGCTTTTT	0,94	2	2,13		Hs.102267	lysyl oxidase
2202	ATTTTCATCAA	0,94	2	2,13	0,46	Hs.101282	Homo sapiens mRNA; cDNA DKFZp434B102 (from clone
2203	TACAGAATGTG	0,47	1	2,13		Hs.99196	ESTs
2204	TTGTATTGTTG	0,47	1	2,13	0,23	Hs.98445	ESTs
2205	GGGCAGAAGGC	0,47	1	2,13	0,23	Hs.9585	ESTs
	TATTGGCCTGG	0,47	1	2,13	0,23	Hs.79572	cathepsin D (lysosomal aspartyl protease)
	TGACTGTAAAA	0,47	1	2,13		Hs.75621	protease inhibitor 1 (anti- elastase), alpha-1-antitr
	ACCTGCTTCCC	0,47	1	2,13		Hs.75458	ribosomal protein L18
	TGATACAGAAA	0,47	1	2,13		Hs.69504	ESTs
2210	AATGAATTCTT	0,47	1	2,13	0,23	Hs.5613	Homo sapiens mRNA, cDNA DKFZp564E2222 (from
2211	ATTTTGTCACT	0,47	1	2,13	0,23	Hs.5459	KIAA1436 protein
2212	CACCCTTTACT	0,47	1	2,13	0,23	Hs.5437	Tax1 (human T-cell leukemia virus type I) binding pr
	CTGGTGGGCCA	0,47		2,13		Hs.5338	carbonic anhydrase XII
	CTTTGAAATAG	0,47		2,13		Hs.50476	ESTs
_	AGTCCTGCTTC	0,47	1	2,13	0,23	Hs.44565	ESTs
	TAACATTGAGA	0,47	1	2,13		Hs.43756	ESTs
2217	CTGAGAAGCGG	0,47	1	2,13	0,23	Hs.41055	ESTs, Highly similar to RL2A_HUMAN 60S RIBOSOMAL

		0.47		0.40			It is a second of the second
2218	TATATATAGAG	0,47	1	2,13	0,23	Hs.34853	inhibitor of DNA binding 4,
0040	CTOTOCTOCAC	0.47		2 42	0.00	Us 22079	dominant negative helix-
2219	GTCTGCTCCAG	0,47	7	2,13	0,23	Hs.32978	proprotein convertase
0000	0000004000T	0.47		0.40	0.00	11- 24040	subtilisin/kexin type 7
	CCCCGAGGCT	0,47	1	2,13		Hs.31019	ESTs
[ 2221]	AATGAAAAATT	0,47	1	2,13	0,23	Hs.30888	cytochrome c oxidase
1 1						"	subunit VIIa polypeptide 2
0000	TTTOTAGTAGT	- 47		0.40	0.00	11- 00040	like
2222	TTTGTACTAAT	0,47	1	2,13	0,23	Hs.29846	Human DNA sequence
	TTOOATTATOA	0.47		0.40	0.00	11- 00050	from clone 717M23 on
	TTCCATTATCA	0,47		2,13		Hs.29356	ESTs
	GTTTAAAAAGC	0,47		2,13		Hs.29032	ESTs
	CAATTTAAAGT	0,47		2,13		Hs.286249	ESTs
	TGTATTTGTAA	0,47		2,13		Hs.286056	ESTs
	GTGGCGTGCCT	0,47	1	2,13		Hs.282652	EST
2228	TAAAATGTGAC	0,47	1	2,13	0,23	Hs.268447	ESTs, Weakly similar to
							ALU8_HUMAN ALU
2229	AACAATAAAAA	0,47	1	2,13	0,23	Hs.268135	ESTs, Weakly similar to
							ALU4_HUMAN ALU
	GAAACACGTAG	0,47		2,13		Hs.26407	ESTs
2231	ACTTTAACAGG	0,47	1	2,13	0,23	Hs.252387	cadherin EGF LAG seven-
							pass G-type receptor 1
	AGCCCCACAAA	0,47	1	2,13		Hs.250570	ESTs
	TGGTGAGATGA	0,47	1	2,13		Hs.224829	ESTs
	GGCACCGCGTG	0,47	1	2,13		Hs.20677	KIAA1303 protein
2235	CAAGGATAAGA	0,47	1	2,13	0,23	Hs.2001	thromboxane A synthase 1
			•			ă.	(platelet, cytochrome
							P450,
	CGCCCCGGCGG	0,47	1	2,13		Hs.196244	ESTs
2237	GAAGAGAAGGT	0,47	1	2,13	0,23	Hs.180455	RAD23 (S. cerevisiae)
				- 1			homolog A
2238	CAGATTTTGTG	0,47	1	2,13	0,23	Hs.177656	calmodulin 1
		1		ĺĺ			(phosphorylase kinase,
		- 45	- 4	2.13			delta)
	ATTAGTTACAA	0,47		2,13		Hs.177635	KIAA1095 protein
	CCCACCACATT	0,47		2,13		Hs.170610	ESTs
2241	CTGCTAACCCA	0,47	1	2,13	0,23	Hs.170310	cat eye syndrome
		-					chromosome region,
0015	07707070	0.15		0.45		11 400075	candidate 1
2242	GTTCTCTGCTT	0,47	1	2,13	0,23	Hs.169078	Human clone A9A2BRB6
0010	TATOTACACT	- 4-				11- 400004	(CAC)n/(GTG)n repeat-
2243	TATGTACAGTT	0,47	1	2,13	0,23	Hs.163001	hypothetical protein
0011	1400000010					11 404074	PRO0907
	AACCCGGGAGT	0,47		2,13		Hs.161974	EST
	GTAACCTCAAA	0,47		2,13		Hs.161930	EST
2246	TTCAATTTCTT	0,47	1	2,13	0,23	Hs.159971	SWI/SNF related, matrix
{						{	associated, actin
00:	400744					11 45050	dependent
2247	AGCTAAAAAA	0,47	1	2,13	0,23	Hs.15953	hypothetical protein
		i í		1		1	FLJ10120

							1.7.4.
	TTATTTAAAAG	0,47		2,13		Hs.158135	KIAA0981 protein
2249	GACTCCACATT	0,47	1	2,13	0,23	Hs.156637	Cas-Br-M (murine)
							ectropic retroviral
							transforming s
2250	AGACAAAATTA	0,47	1	2,13	0,23	Hs.153487	signal transducing adaptor
	)						molecule (SH3 domain
							and
2251	ACTGAGAAGAG	0,47	1	2,13	0,23	Hs.130761	ESTs, Moderately similar
	8						to AF151803_1 CGI-45
						·	protei
2252	GGTGCTGAATA	0,47		2,13		Hs.12489	ESTs
2253	GATTTCTTTGA	0,47	1	2,13	0,23	Hs.119651	glypican 3
2254	TGTCAAAAGAG	0,47	1	2,13	0,23	Hs.118578	Homo sapiens cDNA
							FLJ20053 fis, clone
							COL00809
2255	ACGGGGAGAGT	0,47	1	2,13	0,23	Hs.117582	CGI-43 protein
2256	TGGCTTTATCC	0,47		2,13		Hs.11506	Human clone 23589
							mRNA sequence
2257	GTTGCGTGTCC	0,47	1	2,13	0,23	Hs.108300	NOT3 (negative regulator
					·		of transcription 3, yeast)
2258	TAAACGGCCTC	0,47	· 1	2,13	0,23	Hs.10632	hypothetical protein
		-/-		,			DKFZp762M136
2259	GGCCAGAATGA	0,47	1	2,13	0.23	Hs.102708	DKFZP434A043 protein
	TGAAGCAAAAA	0,47		2,13		Hs.100407	Homo sapiens mRNA;
		-,	Ĭ	_,	-,		cDNA DKFZp564H2416
							(from
2261	TCACAGCTGTG	8,94	19	2,13	2.52	Hs.77054	B-cell translocation gene
		-,		_,			1, anti-proliferative
2262	AGAAAAAAAA	32,99	70	2,12	7.56	Hs.251680	(Manual assignment) not
				_,			unique, contains loricrin
2263	GTGGCGCACGC	5,19	11	2,12	1.65	Hs.135723	glycolipid transfer protein
	TCTGTACACCT	8,97		2,12		Hs.182740	ribosomal protein S11
	AGGGAGGGCC	2,84		2,11		Hs.172153	glutathione peroxidase 3
		,_		_,	,,,,,		(plasma)
2266	AGGACACCGCC	2,37	5	2,11	0.91	Hs.77793	c-src tyrosine kinase
	CGAGGGCACTC	2,37	5	2,11	0.91	Hs.26915	spectrin, beta, non-
	33,10000,010	-,5'		_,''	0,51	0.200 10	lerythrocytic 2
2268	TGCTGCCAGAC	1,9	Λ	2,11	0.77	Hs.79219	RalGDS-like gene;
	1.3010000000	',"	7	_, ' '	) 5,77	10.10219	KIAA0959 protein
2260	TACTGGTTTAT	1,9		2,11	0.77	Hs.30299	IGF-II mRNA-binding
2209	, , , o , o , o , i , i , i	, , ,	-	2,11	) 5,77	13.00233	protein 2
2270	CCCCCACCTAA	7,18	15	2,09	2 04	Hs.77422	proteolipid protein 2
22,0		ا '' ا	13	2,09	2,04	113.11422	(colonic epithelium-
					ļ		enriched)
2271	TGAATGATACG	1,93		2,07	0.76	Hs.278614	protease, serine, 15
	TGAAACTGCAA			2,07		Hs.147189	
	TGAGTCTGCAA	1,93					HYA22 protein
2213	IIGAGICIGGCI	4,83	10	2,07	1,48	Hs.4055	chromosome 21 open
2274	CCACTOTOOTO	1400		0.00	0.00	Un 0045	reading frame 50
2214	GGAGTGTGCTC	14,06	29	2,06	3,36	Hs.9615	myosin regulatory light
l	1	1 1		1	l	l	chain 2, smooth muscle

							isofo
	TCATCGGGCTG	0,97		2,06		Hs.78335	microtubule-associated protein, RP/EB family, member
2276	GAGACCTTGGA	0,97	2	2,06	0,44	Hs.72249	protease-activated receptor 3
2277	CACTGGACGAG	0,97	2	2,06	0,44	Hs.71574	ESTs
2278	TGTACTTATTA	0,97	2	2,06	0,44	Hs.6906	v-ral simian leukemia viral oncogene homolog A (ras
2279	ACCACCCTGTT	0,97	2	2,06	0.44	Hs.4864	KIAA0892 protein
	ATTTCTCATTC	0,97	2	2,06		Hs.36794	D-type cyclin-interacting protein 1
2281	GGACATTTTC	0,97	2	2,06	0,44	Hs.16986	hypothetical protein FLJ11046
	CATCTTAAATG	0,97	2	2,06		Hs.15467	hypothetical protein FLJ20725
	GAATCATTTAT	0,97	2	2,06		Hs.154668	KIAA0391 gene product
	ATCAAATGCAA	4,37	9	2,06	1,36	Hs.79070	v-myc avian myelocytomatosis viral oncogene homolog
	AGCACCAGAAC	1,46		2,05	0,61	Hs.60103	KIAA0690 protein
	TCCTGACCACC	1,46		2,05		Hs.26002	LIM domain binding 1
	CTTATAATCCC	1,46		2,05		Hs.259541	ESTs
	ACCCATCGCCT	1,46	3	2,05		Hs.165428	ESTs
	CCATTGCACTA	1,46	3	2,05	0,61	Hs.115140	ESTs, Weakly similar to ALU1_HUMAN ALU
2290	TACCCCAGAAC	4,39		2,05	1,35	Hs.145320	ESTs
2291	CTCTCACCCTG	8,79	18	2,05	2,27	Hs.75108	ribonuclease/angiogenin inhibitor
	CAAGGGCTTGC	7,35	15	2,04		Hs.156764	RAP1B, member of RAS oncogene family
2293	TTTGCACTTGT	3,43	7	2,04	1,12	Hs.75188	wee1+ (S. pombe) homolog
	GCCAGGAGCTA	3,43		2,04		Hs.18141	ladinin 1
2295	TAAAATACTCC	1,96		2,04	0,75	Hs.8125	Homo sapiens mRNA; cDNA DKFZp586E1521 (from
	TGATTGATTTG	1,96	4	2,04	0,75	Hs.5912	F-box only protein 7
2297	TAAAAACTTTC	1,96		2,04	0,75	Hs.204096	lipophilin B (uteroglobin family member), prostatein
	TCACTGCACTC	7,85	16	2,04	2,06	Hs.261038	ESTs
	ACCAAAGCCCC	4,95	10	2,02		Hs.284281	Human putative ribosomal protein S1 mRNA
	AATAGGGTCAA	3,96	8			Hs.64797	amyloid beta (A4) precursor-like protein 2
	CCTATAGTCCC	5,45		2,02		Hs.140697	ESTs, Weakly similar to unnamed protein product [H.s
	CCTCCCTGCTC	1,49	3	2,01	0,59	Hs.90790	ESTs
2303	ACCTAGCCACT	1,49		2,01		Hs.89463	potassium large

	<del>,</del>						<del>,</del>
							conductance calcium- activated channe
2304	CAGCTCAGCTG	1,49	3	2,01	0,59	Hs.58414	filamin C, gamma (actin-
				_			binding protein-280)
2305	TCAATAAATGT	1,49	3	2,01	0,59	Hs.106747	ESTs, Weakly similar to
							AF217508_1
							uncharacterized b
2306	CACTCACACCC	1,99	4	2,01	0,73	Hs.24447	sigma receptor (SR31747
2207	CTCAAACCTCT	0.40		0.04		11- 070705	binding protein 1)
2307	GTGAAACCTGT	3,49		2,01	1,1	Hs.272795	hypothetical protein FLJ20359
2308	ATCATACCACG	0	2	2,00	28	Hs.97259	ESTs
	AACGGGGCCCT	0	2	2,00		Hs.97203	small inducible cytokine
2000	70.00000001		_	2,00	2,0	113,57205	subfamily A (Cys-Cys), memb
2310	TCCCCGGTCAG	0	2	2,00	2,8	Hs.80562	gelsolin (amyloidosis,
							Finnish type)
	GCGCATCAAAA	0	2	2,00		Hs.59761	ESTs
2312	AATTTCAGGCA	0	2	2,00	2,8	Hs.5476	serine protease inhibitor,
2010	17000107107						Kazal type, 5
	ATCCCACTACT	0	$\frac{2}{2}$	2,00		Hs.3991	ESTs
	ATCCGCTGGGG	0		2,00		Hs.30954	phosphomevalonate kinase
2315	ACTGTGGACTG	0	2	2,00	2,8	Hs.285122	ESTs, Weakly similar to
			,				S53869 laminin beta-2
2216	AGAGAAGAATG	0		2.00	2.0	Hs.2841	chain
	AGGATTGTTTG	0	2	2,00 2,00		Hs.283545	neuromedin U ESTs
	GTAAAGATTTG	0	2	2,00		Hs.278629	ESTs
	ACTATGGATAG	0	2	2,00		Hs.275511	EST
	GTTAGGCACGA	0	2	2,00		Hs.272800	hypothetical protein
				2,00	2,0		FLJ20456
2321	TGCCACCAACA	0	2	2,00	2,8	Hs.271411	beta-site APP-cleaving
							enzyme 2
	AAATGGGAACA	0		2,00		Hs.271226	ESTs
	GGGGTTTGTTT	0	2	2,00		Hs.258455	EST
	ACATAGTCTGA	0		2,00		Hs.25766	ESTs
	TATGCTGAAAT	0		2,00		Hs.255277	ESTs
2326	TTCACTCCATT	0	2	2,00	2,8	Hs.254914	EST, Weakly similar to NICE-1 protein [H.sapiens]
2327	TCTTGGCAACA	0	2	2,00	2,8	Hs.25431	KIAA1219 protein
	CAATAAAATTC	0		2,00		Hs.250236	EST
2329	GGCTTTTGGTT	0	2			Hs.230730	EST, Moderately similar to
01							RLA1_HUMAN 60S
							ACIDIC
	AGTTTGTTTTA	0		2,00		Hs.212570	EST
	TTCCACTGTGA	0		2,00		Hs.198862	fibulin 2
	CAAGTGGGTGT	0		2,00		Hs.187685	ESTs
2333	TGCAGGGACCT	0	2	2,00	2,8	Hs.173043	metastasis-associated 1-
,	l					l	like 1

2334	TACTTCACCCA	0	2	2,00	2,8	Hs.169517	aldehyde dehydrogenase
							5
	CTTTGATTTAT	0	2	2,00		Hs.165590	ribosomal protein S13
2336	тсттстттт	0	2	2,00	2,8	Hs.145211	Homo sapiens mRNA; cDNA DKFZp434K1111 (from
	GTAGCGCCTCC	0	2	2,00	2,8	Hs.143212	cystatin F (leukocystatin)
	CTAGTGGCGGC	0	2	2,00		Hs.142043	ESTs, Weakly similar to ALU8_HUMAN ALU
2339	ACCTCACCTGG	0	2	2,00	2,8	Hs.137585	UDP glycosyltransferase 2 family, polypeptide B11
2340	CTTGTTCAAAA	0	2	2,00	2,8	Hs.137560	ESTs, Moderately similar to AMSH [H.sapiens]
2341	CCTCTTTCCAG	0	2	2,00	2,8	Hs.134615	ESTs
2342	TCTTAGTTCTA	0	2	2,00	2,8	Hs.130729	ESTs
2343	CACCTTATAGT	0	2	2,00	2,8	Hs.117582	CGI-43 protein
2344	AGCTTCTACCA	0	2	2,00	2,8	Hs.11261	small proline-rich protein 2A
2345	GTCAAGCCCAA	0	2	2,00	2,8	Hs.105033	ESTs, Weakly similar to SPR2J protein [M.musculus]
2346	AACTTATCATT	. 0	2	2,00	2,8	Hs.103368	ESTs
2347	ATGTGCTTCCG	1	2	2,00	0,43	Hs.76494	proline arginine-rich end leucine-rich repeat protei
2348	CCTTTTGTCC	1	2	2,00	0,43	Hs.62601	Homo sapiens mRNA; cDNA DKFZp586K1318 (from
2349	CTCTGTTTACA	1	2	2,00	0,43	Hs.5947	mel transforming oncogene (derived from cell line NK
2350	GAAGTGCTGCT	1	2	2,00	0,43	Hs.21812	ESTs
2351	CCATTGCAGTC	1	2	2,00	0,43	Hs.207659	EST
2352	CCACTGCACGC	1	2	2,00	0,43	Hs.202669	thiopurine S- methyltransferase
	GTGGTGTACAC	1	2	2,00		Hs.168102	Human proteinase activated receptor-2 mRNA, 3'UTR
	ATAAATAAATT	1	2	2,00		Hs.16677	hypothetical protein FLJ10506
	TTGATAAATAA	1	2			Hs.139226	replication factor C (activator 1) 2 (40kD)
	GCAAAAGCCCG	1	2	2,00		Hs.109798	G8 protein
i	TAAAATATGGG	0,5	1	2,00		Hs.98401	Homo sapiens mRNA full length insert cDNA clone EURO
	CTTTGGTTTGC	0,5	1	2,00		Hs.8895	ESTs
2359	CCAAAGGAGAA	0,5	1	2,00		Hs.8889	serine hydroxymethyltransferase 1 (soluble)
2360	TGTACCTTTTC	0,5	1	2,00	0,22	Hs.8172	ESTs

2204	TOTTANCTOTT	0.5		2 00	0.33	Un 70025	matrin 3
	TOTTAAGTGTT	0,5		2,00		Hs.78825	matrin 3
	TACAAGCTGAG	0,5	1	,	·	Hs.77508	glutamate dehydrogenase 1
2363	TCTGCGGGTGG	0,5	1	2,00	0,22	Hs.76159	ATPase, H+ transporting, lysosomal (vacuolar proton
2364	TATCAATATTC	0,5	1	2,00	0,22	Hs.7306	secreted frizzled-related protein 1
2265	ATGTTTTGTAA	0,5	1	2,00	0.22	Hs.6853	carbohydrate (N-
2305	MIGHTIGIAM	0,5	ı	2,00	0,22	1 18.0000	acetylglucosamine 6-O)
		1					sulfotransfer
2366	GCTTTTGTTG	0,5	1	2,00	0,22	Hs.6048	FEM-1-like death receptor
							binding protein
2367	CTGGAGACCCA	0,5	1	2,00	0,22	Hs.56729	lymphocyte-specific
2222	TTOACATAAA	<del>-                                    </del>		2.00	0.00	Un 50040	protein 1
	TTCAGAATAAA	0,5	1			Hs.50848	hypothetical protein FLJ20331
2369	ACAATGAAGCA	0,5	1	2,00	0,22	Hs.285665	Homo sapiens mRNA;
		1					cDNA DKFZp564P013
0070	0007070000	<del> \  </del>		0.00	0.00	11- 005505	(from clone
	GCCTGTGGGGT	0,5	1	,		Hs.285565	ESTs
	CAGCAAGGCTT	0,5	1	2,00		Hs.283714	30 kDa protein
	CAAGGCACCAA	0,5		2,00		Hs.28107	ESTs
	TACCAGAGTCC	0,5	1	2,00		Hs.279927	hypothetical protein
2374	ATTTTTGTAAA	0,5	1	2,00	0,22	Hs.27413	adaptor protein containing
							pH domain, PTB domain
0075	TOTOTTAGAGG			2.00	0.00	Un 070750	and
	TCTGTTACACC	0,5	1			Hs.272759	KIAA1457 protein
	ACCOCATOTTC	0,5				Hs.27263	KIAA1458 protein
	AAGAGATGTTC	0,5	1	2,00		Hs.26799	DKFZP564D0764 protein
	TTGGTCAGGGT	0,5				Hs.264381	EST
	TGGAGCTATGA	0,5		2,00		Hs.261655	EST
	AGACTCAGGCC	0,5		2,00		Hs.24305	ESTs Highly similar to
2381	GTTCATCCTTG	0,5	. 1.	2,00	0,22	Hs.236894	ESTs, Highly similar to LRP1_HUMAN LOW-
							DENSITY
	TATCTTGTTGC	0,5	1	2;00	0,22	Hs.23296	ESTs
2383	GTTTGTTTCCT	0,5	1	2,00	0,22	Hs.21143	ESTs, Weakly similar to
					>		KIAA1532 protein
	44040407077			0.00		11- 00-0-	[H.sapiens]
	AAGACACTGTT	0,5		2,00		Hs.20707	hypothetical protein R31240_1
	CCTAGAATCTG	0,5		2,00		Hs.20196	adenylate cyclase 9
	GTCGGGGGAGA	0,5		2,00		Hs.18844	ESTs
	ATGTTGTCAAT	0,5		2,00		Hs.1845	MHC class I region ORF
2388	TTTTCCCTCAG	0,5	1	2,00	0,22	Hs.184242	sterol-C5-desaturase
		]					(fungal ERG3, delta-5-
							desaturas
2389	ACATTTCATTA	0,5	1	2,00	0,22	Hs.18192	Ser/Arg-related nuclear
							matrix protein (plenty of pr
2390	TTAGCCAGGGT	0,5	1	2,00	0,22	Hs.180610	splicing factor

							proline/glutamine rich (polypyrimidi
2391	GGTGGAAAAA	0,5	1	2,00	0,22	Hs.178728	methyl-CpG binding domain protein 3
2392	GGGCAGAATAA	0,5	1	2,00	0,22	Hs.164690	ESTs
2393	GGTGCCCGGCA	0,5	1	2,00	0,22	Hs.163593	ribosomal protein L18a
2394	TAGCTGAGGCA	0,5	1	2,00	0,22	Hs.159557	karyopherin alpha 2 (RAG cohort 1, importin alpha 1)
2395	CGGATTCAATT	0,5	1	2,00	0,22	Hs.14763	ESTs
2396	AAGAAAGGGAA	0,5	1	2,00	0,22	Hs.140908	ESTs
2397	TCAAAAGGGCA	0,5	1	2,00	0,22	Hs.125158	ESTs
2398	CTCTCCTTGCC	0,5	1	2,00	0,22	Hs.118738	KIAA0800 gene product
2399	TTGCAGAGGGG	0,5	1	2,00	0,22	Hs.110373	ESTs
2400	TGTCTGCAGAA	0,5	1	2,00	0,22	Hs.107418	ESTs
2401	AGCTTCCGCTT	0,5	1	2,00	0,22	Hs.106529	CGI-65 protein
2402	CTCACACACAC	0,5	1	2,00	0,22	Hs.104311	novel protein with MAM domain

Tabelle 7:

Nie	T 0	T 1	1 1 2	10	I to to	
Nr.	Tag-Sequenz	relative Expres sionsfr equenz CGAP	relative Expres sionsfr equenz Haut	Quotie nt	Unigene Accessio n Nr. (NCBI)	Genname und Beschreibung (NCBI)
1	GAGAGCTAACT	0,09	19	211,11	73995	Human profilaggrin mRNA, 3' end
2	GCTGCCCTGGG	0,09	15	166,67	286084	Homo sapiens MRIP- 1 protein (MRIP-1), mRNA
3	GGGTCTGAGGG	0,15	23	153,33	291317	nx40e10.s1 Homo sapiens cDNA, 3' end
4	ACCTGGAGGGG	0,18	20	111,11	135188	602625439F1 Homo sapiens cDNA, 5' end
5	AGGGTGGTGAG	0,15	15	100,00	44036	Homo sapiens cDNA: FLJ21725 fis, clone COLF1013
6	CCACCGCCTGC	0,09	4	44,44	198870	Homo sapiens hypothetical protein MGC10526 (MGC10526), mRNA
7	GACGATGTATA	0,09	4	44,44	1837	POU domain, class 3, transcription factor 1, mRNA
8	GGCCACGTAGC	1,82	63	34,62	155597	Homo sapiens D component of complement (adipsin) (DF), mRNA
9	ACCAAGGACAG	0,18	6	33,33	117938	Human DNA sequence from clone RP11-16H23 on chromosome 10. Contains the gene KIAA0204 (HSLK) for a protein kinase, the COL17A1 gene for collagen type XVII alpha 1 (BP180), ESTs and GSSs
10	AGATTTGGGTA	0,09	3	33,33	314296	QV0-LT0014-250200- 129-c05 Homo sapiens cDNA
11	AGGAACTCTGA	0,09	3	33,33	284202	Homo sapiens X- prolyl aminopeptidase
12	ATGCTGGACAC	0,12	4	33,33	143773	EST384086 Homo sapiens cDNA
13	GAGGAGTCCAT	0,09	3	33,33	1432	Homo sapiens protein kinase C substrate

	<del></del>	<del></del>	Т		<del></del>	Leok H (DDKCCH)
						80K-H (PRKCSH), mRNA
14	CCTCTTTGCAT	0	27	27,00	707	keratin 2A (epidermal ichthyosis bullosa of Siemens)
15	GCAGTGCGTGC	0,15	4	26,67	202508	601649719F1 Homo sapiens cDNA, 5' end
16	CAGAGCCTGCA	0,12	3	25,00	131819	Sushi domain (SCR repeat) containing
17	CGCAACTTCAG	0,12	3	25,00	326035	Homo sapiens early growth response 1 (EGR1), mRNA
18	CGTCTGTAATC	0,12	3	25,00	271661	qh68d02.x1 Homo sapiens cDNA, 3' end
19	GCGTCGGTGCA	0,29	7 .	24,14	155597	Homo sapiens D component of complement (adipsin) (DF), mRNA
20	CCCAGTTAAAA	0,21	5	23,81	7844	golgi autoantigen, golgin subfamily b, macrogolgin (with transmembrane signal), 1
21	AAACCCCATCT	0,09	2	22,22	301228	Homo sapiens cDNA FLJ14144 fis, clone MAMMA1002909
22	AACCCTGGAGA	0,09	2	22,22	61796	Human transcription factor ERF-1 mRNA, complete cds
23	AAGATAATAAA	0,09	2	22,22	251440	ht86b11.x1 Homo sapiens cDNA
24	ACACTAGGCAA	0,09	2	22,22	207407	hypothetical protein FLJ21195 similar to protein related to DAC and cerberus
25	AGCCTGGGAAG	0,09	2	22,22	307069	"Homo sapiens mRNA; cDNA DKFZp547N032 (from clone DKFZp547N032)"
26	AGCTGAGTTTG	0,09	2	22,22	288760	Homo sapiens cDNA FLJ12327 fis, clone MAMMA1002140
27	AGGCAATGTTT	0,09	2	22,22	259090	wd43b08.x1 Homo sapiens cDNA, 3' end
28	ATGAAGAAGGA	0,18	4	22,22	2780	Homo sapiens jun D proto-oncogene (JUND), mRNA
29	CAGGAGGACTG	0,09	2	22,22	323833	Homo sapiens syntaphilin (KIAA0374), mRNA

30	CATCCCTTGCC	0,09	2	22,22	146246	601482319F1 Homo sapiens cDNA, 5' end
31	CGCCAGTAATC	0,09	2	22,22	260970	za71f05.x5 Homo sapiens cDNA, 3' end
32	CTCTTGTACTG	0,09	2	22,22	233377	602016005F1 Homo
33	CTTTGTGATGC	0,09	2	22,22	224080	sapiens cDNA, 5' end hx98g05.x1 Homo
34	GAGTGAGCAGA	0,09	2	22,22	130760	sapiens cDNA, 3' end Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 12B (PPP1R12B), transcript variant 2, mRNA
35	GATGTTGATAA	0,09	2	22,22	102243	AV708310 Homo sapiens cDNA, 5' end
36	GCCCACTGCTC	0,09	2	22,22	187617	Homo sapiens cDNA FLJ13941 fis, clone Y79AA1000850
37	GTGAGGCTTTG	0,09	2	22,22	334302	RST29457 Homo sapiens cDNA
38	GTGTTGCACGC	0,09	2	22,22	191341	Homo sapiens cDNA FLJ12161 fis, clone MAMMA1000576
39	TCAAAACCCAA	0,09	2	22,22	282415	AV649184 Homo sapiens cDNA, 3' end
40	TCTCAAAATAA	0,09	2	22,22	298117	UI-H-BW1-anv-g-11- 0-UI.s1 Homo sapiens cDNA, 3' end
41	TCTTCTGAAAA	0,09	2	22,22	322645	"Homo sapiens mRNA; cDNA DKFZp586J101 (from clone DKFZp586J101)"
42	TGCCTGTAAGT	0,09	2	22,22	102325	yv65a03.s1 Homo sapiens cDNA, 3' end
43	TGGGCAGGTGT	0,09	2	22,22	333170	Homo sapiens MAP1 light chain 3-like protein 2 mRNA, complete cds
44	ТСТТТТТСТС	0,09	2	22,22	167406	602624583F1 Homo sapiens cDNA, 5' end /clone=IMAGE:47493 32
45	GGAGAGAAAAG	0,23	5	21,74	158291	Human DNA sequence from clone RP1-233K16 on chromosome 1p36.21-36.33 Contains the gene

	<del></del>					
						KIAA0444, a putative chromodomain helicase DNA binding protein 3 (CHD3), the gene for potassium channel beta 2 subunit (KCNK2), two CpG island, ESTs, STSs and GSSs
46	AGCCTAGGAGT	0,15	3	20,00	37308	yq98e10.s1 Homo sapiens cDNA, 3' end
47	CCAGCGCCAAC	0,7	14	20,00	11950	GPI-anchored metastasis-associated protein homolog
48	CCATCTTGAGG	0,15	3	20,00	110707	H326
49	GCAAAATTCTG	0,15	3	20,00	336728	nk76b08.s1 Homo sapiens cDNA, 3' end /clone=IMAGE:10194 15
50	GCTTTCACCCA	0,26	5	19,23	35841	nuclear factor I/X (CCAAT-binding transcription factor)
51	CCTGTAATCAA	0,21	4	19,05	167596	AV735893 Homo sapiens cDNA, 5' end
52	GTGTGGTGGAG	0,32	6	18,75	177486	amyloid beta (A4) precursor protein (protease nexin-II, Alzheimer disease)
53	CTCGTGGGAAA	0,23	4	17,39	118727	Homo sapiens cDNA FLJ13692 fis, clone PLACE2000103
54	GTGGTGCACAG	0,23	4	17,39	32945	glutamate receptor, metabotropic 1
55	ACGGGGAAAAA	0,12	2	16,67	178357	Homo sapiens cDNA FLJ13657 fis, clone PLACE1011563
56	CAAACTTTAAT	0,12	2	16,67	213069	wd38c05.x1 Homo sapiens cDNA, 3' end
57	CACAGTAGCTC	0,12	2	16,67	269339	EST375707 Homo sapiens cDNA
58	CCCCTGTGCTC	0,12	2	16,67	289077	Homo sapiens cDNA FLJ12768 fis, clone NT2RP2001576, weakly similar to HYPOTHETICAL 62.2 KD PROTEIN C4G8.12C IN CHROMOSOME I
59	CCCGGTTAATT	0,12	2	16,67	325049	no94f01.s1 Homo sapiens cDNA /clone=IMAGE:11144

			1	T		89
60	CGCAACCTCAT	0,12	2	16,67	26557	plakophilin 3
61	CGTTTAAGTCA	0,12	2	16,67	85146	Homo sapiens chromosome 21 derived BAC containing erythroblastosis virus
						oncogene homolog 2 protein (ets-2) gene, complete cds
62	CTAGCCTGGGT	0,12	2	16,67	245025	xd92d10.x1 Homo sapiens cDNA, 3' end
63	GCAGACACCTG	0,12	2	16,67	81234	Homo sapiens mRNA for KIAA0466 protein, partial cds
64	GCAGTGGTGAG	0,12	2	16,67	107003	Homo sapiens enhancer of invasion 10 (HEI10), mRNA
65	GCCAACATAGT	0,12	2	16,67	287722	Homo sapiens cDNA: FLJ23104 fis, clone LNG07676
66	GCCAGTGCCCT	0,12	2	16,67	93183	vasodilator-stimulated phosphoprotein
67	GCGAAAACCCT	0,12	2	16,67	317385	yb56a02.r1 Homo sapiens cDNA, 5' end
68	GGAGAGCAGCA	0,12	2	16,67	177596	Homo sapiens clone PP2464 unknown mRNA
69	GGAGGGAGTTT	0,12	2	16,67	16007	Homo sapiens genomic DNA, chromosome 21q, section 89/105
70.	GTGGGAGGATT	0,18	3	16,67	122309	Homo sapiens mRNA for KIAA1661 protein, partial cds
71	TGCAAAGGCTG	0,12	2	16,67	13801	Homo sapiens mRNA for KIAA1685 protein, partial cds
72	CCTTGGGCACA	0	16	16,00	507	Homo sapiens corneodesmosin (CDSN), mRNA
73	GTCAGTTCCTG	0,38	6	15,79	3796	EphB6
74	GTGTCACGTGC	0,26	4	15,38	336189	"Homo sapiens mRNA; cDNA DKFZp434F1135 (from clone DKFZp434F1135)"
75	AGGAGGTCGCT	0,21	3	14,29	74316	Homo sapiens desmoplakin (DPI, DPII) (DSP), mRNA
76	GTGGTGAGTAC	0,21	3	14,29	182999	yx41g11.s1 Homo

				•		sapiens cDNA, 3' end
77	AGAATGAGATC	0,56	8	14,29	76152	Homo sapiens decorin (DCN), mRNA
78	AAGGCCAGCAA	0,15	2	13,33	25333	Homo sapiens interleukin 1 receptor, type II (IL1R2), mRNA

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## Patentansprüche:

Verfahren zur Identifizierung der in Haut exprimierten Gene bei Menschen 1. oder Tieren in vitro, dadurch gekennzeichnet, daß man-

- a) ein Gemisch von in menschlicher oder tierischer Haut exprimierten genetisch codierten Faktoren aus menschlicher oder tierischer Haut gewinnt und
- b) das in a) gewonnenen Gemisch einer Seriellen Analyse Genexpression (SAGE) unterwirft, und dadurch die in menschlicher oder tierischer Haut exprimierten Gene identifiziert und ihre Expression quantifiziert.
- Verfahren zur Identifizierung der für die Homeostase der Haut bedeutsamen Gene bei Menschen oder Tieren in vitro, dadurch gekennzeichnet, daß man
  - a) ein Gemisch von in menschlicher oder tierischer Haut exprimierten genetisch codierten Faktoren aus menschlicher oder tierischer Haut gewinnt,
  - b) das in a) gewonnenen Gemisch einer Seriellen Analyse Genexpression (SAGE) unterwirft, und dadurch die in menschlicher oder tierischer Haut exprimierten Gene identifiziert und ihre Expression quantifiziert und
  - c) die Analysergebnisse aus b) mit Expressionsmustern anderer Gewebe vergleicht und so die Gene identifiziert, die in Haut und anderen Geweben unterschiedlich stark (differentiell) exprimiert werden.
- 3. Verfahren zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, dadurch gekennzeichnet, daß man
  - a) ein Gemisch von Proteinen, mRNA-Molekülen oder Fragmenten von Proteinen oder mRNA-Molekülen aus menschlicher oder tierischer Haut gewinnt,
  - b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder

Fragmente von Proteinen oder mRNA-Molekülen untersucht, die mittels Serieller Analyse der Genexpression (SAGE) als in Haut und anderen Geweben unterschiedlich stark (differentiell) exprimiert identifiziert werden.

- c) die Untersuchungsergebnisse aus b) mit den mittels Serieller Analyse der Genexpression (SAGE) identifizierten Expressionsmustern vergleicht und
- d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut stärker exprimiert werden als in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben stärker exprimiert werden als in Haut.

## 4. Verfahren nach Anspruch 3, dadurch gekennzeichnet, daß man

in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 1 bis 5 und 7 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens doppelt so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Molekülen oder Fragmente von Proteinen oder mRNA-Molekülen

enthält, die in anderen Geweben mindestens doppelt so stark exprimiert werden wie in Haut.

5. Verfahren nach Anspruch 3 oder 4, dadurch gekennzeichnet, daß man in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 2 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden, in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 2 bis 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 5-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine. mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 5-fach so stark exprimiert

 Verfahren nach einem der Ansprüche 3 bis 5, dadurch gekennzeichnet, daß man

werden wie in Haut.

in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 3 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 3 bis 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 10-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Molekülen oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 10-fach so stark exprimiert werden wie in Haut.

## Verfahren nach einem der Ansprüche 3 bis 6, dadurch gekennzeichnet, daß man

in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 4 und 5 in Spalte 7 durch ihre UniGene-Accession-Number definiert werden,

in Schritt c) die Untersuchungsergebnisse aus b) mit den in den Tabellen 4 in den Spalten und 3 und angegebenen relativen Expressionsfrequenzen sowie Spalte 5 angegebenen den in Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 20-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Molekülen oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 20-fach so stark exprimiert werden wie in Haut.

8. Verfahren nach einem der Ansprüche 3 bis 7, dadurch gekennzeichnet, daß man

in Schritt b) das gewonnene Gemisch auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in Tabelle 5 in Spalte 7 durch ihre UniGene-Accession-Number definiert werden.

in Schritt c) die Untersuchungsergebnisse aus b) mit den in Tabelle 5 in den Spalten 3 und 4 angegebenen relativen Expressionsfrequenzen sowie den in Spalte 5 angegebenen Expressionsquotienten vergleicht und

in Schritt d) das in b) untersuchte Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in Haut mindestens 100-fach so stark exprimiert werden wie in anderen Geweben, oder das in b) untersuchte Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn es überwiegend Proteine, mRNA-Molekülen oder Fragmente von Proteinen oder mRNA-Molekülen enthält, die in anderen Geweben mindestens 100-fach so stark exprimiert werden wie in Haut.

- Verfahren zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, dadurch gekennzeichnet, daß man
  - a) ein Gemisch von Proteinen, mRNA-Molekülen oder Fragmenten von Proteinen oder mRNA-Molekülen aus menschlicher oder tierischer Haut gewinnt,
  - b) in dem gewonnenen Gemisch mindestens zwei der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen quantifiziert, die mittels eines Verfahrens nach Anspruch 2 als für die Homeostase der Haut bedeutsam identifiziert werden.
  - c) die Expressionsverhältnisse der mindestens zwei Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen zueinander bestimmt,

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- d) die Expressionsverhältnisse aus c) mit den Expressionsverhältnissen vergleicht, die für die in b) quantifizierten Moleküle typischerweise in homeostatischer vorliegen, insbesondere Haut Expressionsverhältnissen, die sich aus Tabelle 6, Spalte 3 bzw. aus den Tabellen 1 bis 5, Spalte 4 ergeben, und
- e) das in a) gewonnene Gemisch gesunder bzw. in Homeostase befindlicher Haut zuordnet, wenn die Expressionsverhältnisse der untersuchten Haut Expressionsverhältnissen in Homeostase befindlicher Haut entsprechen, oder das in a) gewonnene Gemisch kranker bzw. in gestörter Homeostase befindlicher Haut zuordnet, wenn die Expressionsverhältnisse der untersuchten Haut von den Expressionsverhältnissen in Homeostase befindlicher Haut abweichen.
- 10. Verfahren nach einem der Ansprüche 1 bis 9, dadurch gekennzeichnet, daß man in Schritt a) das Gemisch aus einer Hautprobe, insbesondere aus einer Vollhautprobe oder aus einer Epidermisprobe gewinnt.
- Verfahren nach einem der Ansprüche 3 bis 9, dadurch gekennzeichnet, daß man in Schritt a) das Gemisch mittels Mikrodialyse gewinnt.
- 12. Verfahren nach einem der Ansprüche 3 bis 8, 10 und 11, dadurch gekennzeichnet, daß man die Untersuchung in Schritt b) auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der Proteine oder Proteinfragmente mittels einer Methode durchführt, die ausgewählt ist unter
  - Ein- oder zweidimensionaler Gelelektrophorese
  - Affinitätschromatographie
  - Protein-Protein-Komplexierung in Lösung
  - Massenspektrometrie, insbesondere Matrix Assistierter Laser Desorptions Ionisation (MALDI) und insbesondere
  - Einsatz von Proteinchips,

oder mittels geeigneter Kombinationen dieser Methoden.

- 13. Verfahren nach einem der Ansprüche 9 bis 11, dadurch gekennzeichnet, daß man in Schritt b) die Quantifizierung mindestens zweier Proteine oder Proteinfragmente mittels einer Methode durchführt, die ausgewählt ist unter
  - Ein- oder zweidimensionaler Gelelektrophorese
  - Affinitätschromatographie
  - Protein-Protein-Komplexierung in Lösung
  - Massenspektrometrie, insbesondere Matrix Assistierter Laser Desorptions Ionisation (MALDI) und insbesondere
  - Einsatz von Proteinchips, oder mittels geeigneter Kombinationen dieser Methoden.
- 14. Verfahren nach einem der Ansprüche 3 bis 8, 10 und 11, dadurch gekennzeichnet, daß man die Untersuchung in Schritt b) auf das Vorhandensein und gegebenenfalls die Menge von mindestens einem der mRNA-Moleküle oder mRNA-Molekülfragmente mittels einer Methode durchführt, die ausgewählt ist unter
  - Northern Blots,
  - Reverse Transkriptase Polymerasekettenreaktion (RT-PCR),
  - RNase-Schutzexperimente,
  - Dot-Blots,
  - CDNA-Sequenzierung,
  - Klon-Hybridisierung,
  - Differential Display,
  - Subtraktive Hybridisierung,
  - · cDNA-Fragment-Fingerprinting,
  - Total Gene Expression Analysis (TOGA)
  - Serielle Analyse der Genexpression (SAGE) und insbesondere
  - Einsatz von Nukleinsäurechips,

oder mittels geeigneter Kombinationen dieser Methoden.

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- 15. Verfahren nach einem der Ansprüche 9 bis 11, dadurch gekennzeichnet, daß man in Schritt b) die Quantifizierung mindestens zweier mRNA-Moleküle oder mRNA-Molekülfragmente mittels einer Methode durchführt, die ausgewählt ist
  - Northern Blots,
  - Reverse Transkriptase Polymerasekettenreaktion (RT-PCR),
  - RNase-Schutzexperimente,
  - Dot-Blots,

unter

- CDNA-Sequenzierung,
- Klon-Hybridisierung,
- Differential Display,
- Subtraktive Hybridisierung,
- cDNA-Fragment-Fingerprinting,
- Total Gene Expression Analysis (TOGA)
- Serielle Analyse der Genexpression (SAGE) und insbesondere
- Einsatz von Nukleinsäurechips,

oder mittels geeigneter Kombinationen dieser Methoden.

- 16. Verfahren nach einem der Ansprüche 3 bis 8, 10, 11, 12 und 14, dadurch gekennzeichnet, daß man in Schritt b) auf das Vorhandensein und gegebenenfalls die Menge von 1 bis etwa 5000, bevorzugt 1 bis etwa 1000, insbesondere etwa 10 bis etwa 500, vorzugsweise etwa 10 bis etwa 250, besonders bevorzugt etwa 10 bis etwa 100 und ganz besonders bevorzugt etwa 10 bis etwa 100 und ganz besonders bevorzugt etwa 10 bis etwa 50 der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen untersucht, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden.
- 17. Verfahren nach einem der Ansprüche 9 bis 11, 13 und 15, dadurch gekennzeichnet, daß man in Schritt b) 1 bis etwa 5000, bevorzugt 1 bis etwa

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1000, insbesondere etwa 10 bis etwa 500, vorzugsweise etwa 10 bis etwa 250, besonders bevorzugt etwa 10 bis etwa 100 und ganz besonders bevorzugt etwa 10 bis etwa 50 der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen quantifiziert, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden.

- 18. Test-Kit zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, umfassend Mittel zur Durchführung der Verfahren nach einem der Ansprüche 3 bis 17.
- Biochip zur Bestimmung der Homeostase der Haut bei Menschen oder Tieren in vitro, umfassend
  - einen Träger und
  - auf diesem immobilisierte Sonden, die zur spezifischen Bindung an mindestens eines der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen befähigt sind, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden.
- 20. Biochip nach Anspruch 19, umfassend 1 bis etwa 5000, bevorzugt 1 bis etwa 1000, insbesondere etwa 10 bis etwa 500, vorzugsweise etwa 10 bis etwa 250, besonders bevorzugt etwa 10 bis etwa 100 und ganz besonders bevorzugt etwa 10 bis etwa 50 voneinander verschiedene Sonden.
- 21. Biochip nach Anspruch 19 oder 20, umfassend Nukleinsäuresonden, insbesondere RNA- oder PNA-Sonden, besonders bevorzugt DNA-Sonden.
- 22. Biochip nach Anspruch 21, umfassend Sonden mit einer Länge von etwa 10 bis etwa 1000, insbesondere etwa 10 bis etwa 800, vorzugsweise etwa 100 bis etwa 600, besonders bevorzugt etwa 200 bis etwa 400 Nukleotiden.

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- 23. Biochip nach Anspruch 19 oder 20, umfassend Peptid- oder Proteinsonden, insbesondere Antikörper.
- 24. Verwendung der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden, als Marker für die Homeostase der Haut bei Menschen oder Tieren.
- 25. Testverfahren zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom, in vitro, dadurch gekennzeichnet, daß man
  - a) den Hautstatus durch ein Verfahren nach einem der Ansprüche 3 bis 17, oder mittels eines Test-Kits nach Anspruch 18, oder mittels eines Biochips nach einem der Ansprüche 19 bis 23 bestimmt,
  - b) einen Wirkstoff zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut einmal oder mehrmals auf die Haut aufbringt,
  - c) erneut den Hautstatus durch ein Verfahren nach einem der Ansprüche 3 bis 17, oder mittels eines Test-Kits nach Anspruch 18, oder mittels eines Biochips nach einem der Ansprüche 19 bis 23 bestimmt, und
  - d) die Wirksamkeit des Wirkstoffs durch den Vergleich der Ergebnisse aus a) und c) bestimmt.
- 26. Test-Kit zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut in vitro, umfassend Mittel zur Durchführung des Verfahrens nach Anspruch 25.

- 27. Verwendung der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden, zum Nachweis der Wirksamkeit von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkom.
- 28. Screening-Verfahren zur Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkomin vitro, dadurch gekennzeichnet, daß man
  - a) den Hautstatus durch ein Verfahren nach einem der Ansprüche 3 bis 17, oder mittels eines Test-Kits nach Anspruch 18, oder mittels eines Biochips nach einem der Ansprüche 19 bis 23 bestimmt,
  - b) einen potentiellen Wirkstoff zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut einmal oder mehrmals auf die Haut aufbringt,
  - c) erneut den Hautstatus durch ein Verfahren nach einem der Ansprüche 3 bis 17, oder mittels eines Test-Kits nach Anspruch 18, oder mittels eines Biochips nach einem der Ansprüche 19 bis 23 bestimmt, und
  - d) wirksame Wirkstoffe durch den Vergleich der Ergebnisse aus a) und c) identifiziert.
- 29. Verwendung der Proteine, mRNA-Moleküle oder Fragmente von Proteinen oder mRNA-Molekülen, die in den Tabellen 1 bis 5 in Spalte 7 und in Tabelle 7 in Spalte 6 durch ihre UniGene-Accession-Number definiert werden, zur

Identifikation von kosmetischen oder pharmazeutischen Wirkstoffen zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkomin.

- 30. Verfahren zur Herstellung einer kosmetischen oder pharmazeutischen Zubereitung zur Aufrechterhaltung oder Förderung der Homeostase der Haut bzw. zur Behandlung pathologischer Zustände der Haut, wie Neurodermitis, Sonnenbrand, Psoriasis, Sklerodermie, Ichtyosis, atopische Dermatitis, Akne, Seborrhoe, Lupus erythematodes, Rosacea, Melanoma, Basalioma, Hautkarzinom, Hautsarkomin, dadurch gekennzeichnet, daß man
  - a) wirksame Wirkstoffe mit Hilfe des Verfahrens nach Anspruch 28, oder der Verwendung nach Anspruch 29 bestimmt und
  - b) als wirksam befundene Wirkstoffe mit kosmetisch und pharmakologisch geeigneten und verträglichen Trägern vermischt.

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SEQUENCE LISTING - SEQUENZPROTOKOLL - enthaltend die Sequenzen aus Tabelle 6 der nachfolgend genannten Patentanmeldung

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